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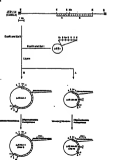
INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCD

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(21) International Application Number: PCT/US j22) International Filing Date: 14 March 1990 (20) Peterly date: 17 March 1990 (17.03.89) (327,05) 17 March 1989 (17.03.89) (30) Peters Application or Grassi (43) Related by Cominantion 17 March 1990 (71) Applicant for all designated States except US: Policy To Extend USA ND COMPANY 1007 March 1990 (17.01.89) (72) Investors; AND COMPANY 1007 March 1990 (17.01.89) (73) Investors; Applicants (for US only) : HERSHEY 1993 (17.01.89) (73) Investors; Applicants (for US only) : HERSHEY 1993 (17.01.89) (73) Investors; Applicants (for US only) : HERSHEY 1993 (17.01.89) (73) Investors; All Company (17.01.89) (74) Investors; All Company (17.01.89) (75) Investors; All Company (17.01.89) (75) Investors; All Company (17.01.89) (75) Investors (17.01.89) (75	(14.03.: (14.03.: (17.03.: E.I. I. US/U: S). Howarder, I US]; 3 (18.); 3 (19.); 3 (1	90) US (P) (P) (P) (P) (P) (P) (P) (P) (P) (P)	(74) Agents: MORRISSEY, Bruce, W. et al.; E.I. du Pont de Nemours and Company, 1007 Market Street, Wilnington, De 1996 (US). (81) Designated States: AT. (European patent), AU, B.B. Be (European patent), Br. (OAPT patent), B.B. (DAPT patent), B.R. (DA, CP (OAPT patent), D.B. (European patent), B.F. (European patent), C. (European patent), D.F. (European patent), G. (OAPT patent), D.F. (European patent), G.F. (UA) (CAPT patent), D.F. (European patent), G.F. (UA) (CAPT patent), M.F. (UR) (CAPT patent), U.S. (UR) (CAPT	
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(54) Title: EXTERNAL REGULATION OF GENE EXPRESSION

(57) Abstract

The preparation and use of nucleic aid promoter impanets derived from several genes from core, petunia and tobacco which are highly responsive to a number of substituted beargeneralifonamides and related compounds are described. These promoter fragments are useful in centaing recombinant DNA constructions comprising nuclearing the company desired gene product operably linked to such promise product operably linked to such promise product under external chemical control in various studies of tender plants and bring the expression of the gene product under external chemical control in various situates of monocorpical control and discoprised control plants.



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TITLE

5 CROSS-REFERENCE TO RELATED APPLICIATION
This application is a continuation-in-part of

U.S. Serial No. 07/327,205, filed March 17, 1989.

FIELD OF THE INVENTION

This invention relates to the preparation and use of nucleic acid promoter fragments derived from several genes from corn, petunia and tobacco which are highly responsive to a number of substituted benzenesulfonamides and related compounds. Chimeric genes consisting of nucleic acid sequences encoding a desired gene product operably linked to one of these promoter fragments in recombinant DNA constructions may be made. Transformation of plants with such constructions will result in new plants in which the

expression of the product encoded by such chimeric genes can be controlled by the application of a suitable inducing chemical.

BACKGROUND OF THE INVENTION

The ability to externally control the expression of selected genes and thereby their gene

products in field-grown plants by the application of appropriate chemical substances in the field can provide important agronomic and foodstuff benefits. This control is especially desirable for the regulation of genes that might be placed into

- 30 transgenic plants and has many applications including (1) prolonging or extending the accumulation of desirable nutritional food reserve in seeds, roots, or tubers, (2) producing and accumulating products in plant tissues at a defined time in the
- developmental cycle such that these products are convenient for harvest and/or isolation, and (3)

	initiating the expression of a pest-specific toxin a
	the site of pathogen attack. The latter example may
5	provide a means of avoiding contamination of the
	ultimate food product with the toxic agent as well a
	minimizing the development of resistance in the pest
	population by selective, tissue specific, rather than
	constitutive expression of the toxic agent. These
10	and other benefits have been unattainable to date
	since a practical means to bring known plant genes
	under external control in the field has not been
	available.
	In eukaryotic systems, the expression of genes
15	is directed by a region of DNA called the promoter.
	In general, the promoter is considered to be that
	portion of DNA in a gene upstream from the coding
	region that contains the site for the initiation of
	transcription. The promoter region also comprises
20	other elements that act to regulate gene expression.
	These include the "TATA box" at approximately 30 bp
	(-30) 5' relative to the transcription start site an
	often a "CAAT box" at -75 bp. Other regulatory
	elements that may be present in the promoter are
25	those that affect gene expression in response to
	environmental stimuli, such as light, nutrient
	availability, heat, anaerobioisis, the presence of
	heavy metals, and so forth. Other DNA sequences
	contained within the promoter may affect the
30	developmental timing or tissue specificity of gene
	expression. In addition, enhancer-like sequences
	that act to increase overall expression of nearby
	genes in a manner that is independent of position or
	orientation have been described in a number of
35	eukaryotic systems. Homologs of these enhancer-like
	sequences have been described for plants as well.

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The vast diversity of promoter function in eukaryotic systems therefore provides the opportunity to isolate

promoters with relatively stringent requirements for their transcriptional activation which may be useful in regulating the timely expression of gene products in transgenic plants.

While current technology exists to transform plants with the genes encoding selected products, the 10 expression of these genes is either continuous throughout the life cycle (controlled by a constitutive promoter), or regulated by the developmentally timed program of maturation inherent 15 in each organ/tissue/cell (stage or tissue specific

promoters) in which the gene product is destined to be expressed. Continuous expression precludes controlled production of a gene product at particular stages of the life cycle, in specific tissues or in

response to environmentally unpredictable events. In addition, such constitutive expression could place a major penalty on yield, due to greatly increased energy demands accompanying prolonged high level synthesis of a single gene product. Tissue or stage 25 specific expression, although valuable for the temporal and spatial accumulation of products, is

under the variable timing of the developmental program of each plant. The practical use of promoters from these types of genes would therefore necessitate the isolation of a multitude of stageand tissue-specific promoters for all crop species of interest.

Ideally, one would prefer to externally control the expression of a gene product in transgenic plants by application of an inducing signal that stimulates expression of the desired gene in any tissue(s) at

any time in the plant's life cycle. This regulation would be accomplished by controlling the expression

5 of a structural gene encoding the desired product with a promoter that is highly responsive to application of the inducing signal. The proposed inducer/promoter combination should be functional in a wide variety of plant species, with the inducer having no effect on the normal plant growth, development or morphology. Chemicals that fit the

development or morphology. Chemicals that fit the above criteria for regulating gene expression in plants would be of great utility in the field, as their use would be compatible with current

agricultural practices. For instance, application of a chemical inducer could be easily accomplished using equipment currently in use by most plant growers.

Ideally, a chemical/chemically responsive promoter

combination could be made functional at any stage or in any tissue of a transformable plant to control the expression of any desired gene product.

There are inducer/promoter combinations that have been shown to regulate the expression of foreign genes in both bacterial and animal systems. Many of

25 the inducible bacterial systems are based on the use of promoters that respond to metabolites or metabolite analogs that normally regulate bacterial growth. Addition of an appropriate metabolite to the media of active growing bacterial cultures trans-

30 formed with genes driven by promoters that are responsive to these metabolites results in expression of the desired product. Examples of such inducer/promoter combinations include

3-β-indoylacrylic acid/Trp promoter, IPTG/lac
35 promoter, phosphate/phosphate starvation inducible

35 promoter, phosphate/phosphate starvation inducible promot r, and L-arabinose/ara B promoter

combinations. Similarly, heavy metal/metallothionine promoter, and heat/heat shock promoter combinations

have been used in animal cell culture systems to control the expression of foreign genes.

There are a number of inducer/promoter combinations derived from plant genes that are known. Activation of many of these promoters is regulated by environmental factors such as light, heat shock and anaerobiosis. The promoters of these inducible genes have been extensively analyzed [c.f.,

Kuhlemeier et al., Ann. Rev. Plant Physiol., 38:221-257 (1987)]. However, the use of environmental inducers for regulating foreign genes

is impractical since the inducing signal(i.e., light, temperature and O₂ levels) are not easily or practically controllable under conditions of normal agranomic practices. Other plant dense have been

described that are induced by oligosaccharides, such as those generated during pathogen infection and/or wounding. Examples include the induction of phenylalanine ammonia lyase and chalcone synthase by glucan elicitors in soybean [Ebel, J., et al., Arch.

25 Biochem. Biophys. 232, 240-248 1984] and induction of a wound-inducible inhibitor gene in potato [Cleveland, T.E. et al., Plant Mol. Biol. 8, 199-208 1987]. Again, the promoters of these inducible genes lack utility in regulating the expression of foreign genes in transformed plants due to either lack of a

practical method of induction (wounding) or the deleterious effects that result from diverting metabolic energy from plant growth to large scale synthesis of products designed to combat pathogen 35 attack (oliogasccharide inducers).

A large number of chemicals, both natural products and synthetic compounds, have potential use

in controlling gene expression in plants. However, any chemical that may be useful as an inducer of gene

- 5 expression in the field must minimally be environmentally safe, have little or no effect on the normal growth, morphology and development of plants, and be easily used under conditions of normal agronomic practice.
- A number of natural products are known that affect gene expression. These are mainly naturally occurring plant growth regulators such as the auxins, cytokinins, gibberellic acid, ethylene and abscisic acid [c.f., Davies, F. (Ed.) Plant Hormones and Their
- Roles In Plant Growth and Development, Martinus Nijhoff Fubl. 1987], while other chemicals have equally dramatic effects such as salicylic acid [Hooft Vanhuijsduijnen et al., J. Gen. Virol.]
 - 67:235-2143 1986]. When the growth regulators
- 20 described above are applied to various plants or plant derived cells/tissues/organs, a change in the metabolism is observed that has been shown to be due, at least in part, to new gene expression. Some products of these genes as well as the genes
- 25 themselves have been isolated and characterized. However, since the chemicals that induce these genes normally function in regulating the growth and development of plants, they cannot be candidates for inducers of recombinant, chemically inducible genes
- 30 in transgenic plants. This lack of utility is a direct result of undesirable pleiotropic effects that would arise from the undesired co-activation of the plant's endogenous hormone sensitive developmental programs along with th desired recombinant gene.
- 35 For example, activation of a foreign gene by abscisic acid in developing plants would induce many

undesirable hormone effects including negative effects on plant metaholism [Milberrow, B.V. An Rev.

- 5 Plant Physiol. 25, 259-307 1974], a sharp decline in growth rate, an induction of stomatal closure, and premature abscission of young leaves and fruits. Other phytohormones have similar negative effects on plant growth and development that preclude their use
- plant growth and development that preclude their use
 in regulating the expression of foreign genes in
 transformed plants. A more general review of
 phytohormone effects on vegetative plants including
 ABA, ethylene, cytokinins, and auxins, is presented
 in Phytohormones and Related Compounds: A
- 15 Comprehensive Treatise Vols I and II, Letham, D.S., Goodwin, P.S., and Higgins, T.G.V. eds. Elsevier/North Holland (1978).

Among the potentially attractive chemical candidates that may have utility in regulating gene

- 20 expression in transgenic plants is the group of compounds collectively called herbicide antidotes or safeners. Safeners are functionally defined as chemicals that have the ability to increase the tolerance of a crop plant to the toxic effects of
- 25 herbicides when the plant is treated with the safener. It now appears that the safening action of these compounds is related to their ability to increase the metabolism of the herbicide in safener-treated plants [Sweetser, P. B., Proceedings of the 1985 British Crop Protection Society
 - Conference-Weeds. 3:1147-1153 1985]. For example, treatment of maire and other cereal crops with safeners such as the dichloroacetamides increases their tolerance toward several groups of herbicides

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[Lay, M.M., and Casida, J.E. Pest. Biochem. Physiol. 6:442-456 1976, Parker, C. Pesticide Science 14:533-536 1983]. More specifically, N,N-diallyl-2.2-dichloroacetamide safening is correlated with an increased level of glutathione-S-transferases (GSTs). a family of enzymes known to detoxify several major classes of pre-emergent, selective herbicides by conjugating them with glutathione [Mozer et al., Biochemistry 22:1068-1072 1983]. This increase in GST activity is correlated with an increased steady-state level of GST mRNA in treated plants, as shown by the work of Wiegand et al [Wiegand, R. et al., Plant Mol. Biol., 7:235-243 1986]. Thus safener treatment of selected plants can increase the steady state level of a gene product without having significant effects on growth and morphology. It has been shown that changes in the rate of metabolic detoxification of sulfonvlurea herbicides in corn plants are induced by treatment with a variety of safeners [Sweetser, P. B., Proceedings of the 1985 British Crop Protection Society Conference. Weeds 3:1147-1153 1985]. The result of this accelerated metabolic detoxification is increased herbicide tolerance in safener-treated plants. For example. 2 to 5 fold increases in the metabolism rates of chlorsulfuron and metsulfuron methyl have been observed in wheat and corn following application of the antidotes napthalic anhydride, N,N-dially1-2,2-dichloroacetamide, or cyometrinil. This observed increase in sulfonylurea herbicide metabolism occurs within hours following antidote treatment. In addition, the safening activity of the

chemicals is not seen if plants are treated with the protein synthesis inhibitor cycloheximide prior to WO 90/11361 PCT/US90/01210

safener treatment, indicating that the increase in herbicide metabolism is dependant on de novo protein synthesis. This requirement for new protein 5 synthesis indicates that safener treatment may activate the transcription of specific nuclear genes. and that a safener/safener-induced gene promoter combination may exist that will have utility in 10 regulating the expression of foreign genes introduced into transgenic plants. To date, however, there has been no reported example of an inducible expression system for transgenic plants based on activation of safener-responsive promoter/structural gene 15 recombinant DNA construction by the external application of a safener or safener like compound. Indeed, no system with real utility for externally regulating the expression of a desired gene in transgenic plants that is compatible with current 20 agronomic practices is known. The instant invention focuses on DNA promoter fragments derived from several plant species which are inducible by herbicide safeners of cereal crops. These promoters have been used to develop a safener/safener inducible gene system for controlling 25 the expression of foreign genes in transformed plants. This system has utility for externally regulating the expression of desired genes in transgenic plants in a grower's field. Its advantages include the high level of activity shown 30 by several of these promoters in response to application of an appropriate inducing chemical, the apparent expression of these promoters in all plant tissues tested to date, and the absence of pleiotr pic effects generated by treatment of plants 35 with these chemicals.

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Ebert et al., [Ebert et al., Proc. Natl. Acad. Sci. (USA) 84:5745-5749 1987], discloses studies of 5 the active fragment of DNA containing the nopaline synthase promoter. This promoter is constitutive rather than inducible, and while of bacterial origin. operates in a wide range of plant tissues. A construction was made so that the promoter controlled 10 the expression of the reporter gene chloramphenicol acetyl transferase (CAT). The authors reported that a fragment of 33 bp (-97 to -130) of DNA was sufficient to promote expression of the CAT gene. They reported further that the presence of two copies 15 of the fragment tripled the expression of the CAT gene. These results from stably transformed tobacco tissue were repeatable in a transient assay using tobacco protoplasts. Comparison of the level of CAT activity obtained when gene expression was controlled 20 by the 33 bp fragment in both the transient expression and stably transformed tobacco protoplasts and tissues resulted in some differences. authors nevertheless indicated their belief that such transient assays are valuable for studies of promoter

expression.

Studies of the anaerobic induction of the maize alcohol dehydrogenase (Adh I) gene by electroporating gene fragments of Adh1 into maize protoplasts from suspension culture cells have been performed [Howard, et al., Planta, 170:535-540 (1987). Transformed protoplasts were subjected to reduced oxygen levels

sequences in stable transformation systems. Operable linkage of the nopeline synthase promoter to a structural gene, however results in constituitive expression of the gene product in transformed plants precluding its use in externally controlling gene

		facilitate
	5	facilitate measurement of anaerobiosis-induced Adhl
	5	gene expression, the 5' promoter or regulatory
r		fragment of the native Adhl gene (1096 base pairs)
		was functionally linked to a CAT gene. Their results
		demonstrated the normal anearobic regulation of the
		inducible Adhl promoter/CAT gene from a monocot maize
	10	gene (i.e., Adhl) in protoplasts derived from a
		homologous cell culture system. They also showed
		that the Adhl promoter fragment, without the coding
		and 3' regions of the Adhl gene, is sufficient for
		anaerobic induction of a foreign coding region in
	15	maize protoplasts.
		Other researchers [Lee et al., Plant Physiology
		85: 327-330 1987], have further defined the size of
		the DNA fragment responsible for anaerobic induction
		of the maize Adhl gene. These researchers
	20	transformed maize protoplasts with a recombinant gene
		consisting of a CAT coding region under the control
		of the Adhl promoter and measured the production of
		CAT 24 hours later. By modifying the length of the
		promoter fragment used in the construction, Lee et
	25	al. determined that 146 bp 5' to the transcription
		start site were sufficient to place the expression of
		CAT under anaerobic induction. However, the
		expression of CAT was increased 5% or 8% by the
		addition of 266 or 955 bp, respectively, of
	30	contiguous 5' promoter sequences.
		Walker et al., [Walker et al., Proc. Natl.
		Acad. Sci. (USA) 84:6634 6639 10073
,		Acad. Sci. (USA) 84:6624-6628 1987], continued the
•		studies of the DNA sequences in the promoter region
	35	of the maize Adhl gene required for aneorbically
	• •	induced gene expression in a transient assay. They

determined that control of anaerobic induction of

35 Ellis et al.

	gene expression resided in two sequences from the
	promoter: those being the sequence between -133 and
5	-124 bp and the sequence between -113 and -99 by (5
	to the transcription start site). Both sequences are
	necessary for induction. Attachment of the full 40
	bp element to an unrelated viral promoter conferred
	anaerobic regulation to the chimeric promoter.
10	Others have shown that extremely low levels of
	CAT gene expression could be observed under
	appropriate anaerobic conditions when the DNA
	fragment between base pairs -1094 and +106 bp of the
	maize Adhl gene was used to regulate CAT gene
15	expression in stably transformed tobacco cells,
	[Ellis et al., EMBO Journal 6:11-16 1987]. In fact,
	only CAT messenger RNA was detected. However,
	promoter elements from the octopine synthase gene of
	bacteria, or those from the Cauliflower Mosaic Virus
20	(CaMV) linked 5' to the Adhl promoter, stimulated the
	expression of the CAT gene and permitted detection of
	CAT after anaerobic induction. The fragment of DNA
	consisting of 247 bp obtained adjacent and 5' to the
	transcription start site of the structural gene for
25	Adhl, was sufficient to put the expression of the CAT
	gene under anaerobic control. Therefore, anaerobic
	control by the 247 bp fragment of DNA was maintained
	even when the octopine synthase and CaMV 358
30	promoters, which are constitutive promoters, were
30	present. The region of the Adhl promoter responsible
	for anaerobic induction demonstrated in transient
	assays by Howard et al., Lee et al., and Walker et
	al. were similar and identical to the region showing
	anaerobic induction in stably transformed plants by

Patents have been issued to animal and microbial systems in which the expression of selected 5 gene sequences have been induced by chemicals that interact with certain regulatory sequences. U.S. Patent 4.579,821 issued to Palmiter and Brinster discloses the isolation of promoter/regulator sequences of the mouse metallothionein-I gene and its 10 use to control the expression of selected DNA sequences operably linked to the promoter by exposure to heavy metal ions or steroid hormones. The expression of thymidine kinase fused to the metallothionein-I promoter was obtained in 15 differentiated cells of adult mice upon administration of cadmium or dexamethasone. U.S. Patent 4,703,005 issued to Nakata and Shinagaua discloses the isolation of a gene for phosphatebinding protein (phoS) to which was fused a foreign 20 gene 3' to phoS. The foreign gene is controlled by phosphate in the culture medium. None of these inventions, though has any potential utility for use

with plants in the field. The heavy metal ions that activate the metallothionein promoter are both toxic to plants and would pose an extreme environmental 25 hazard in the field. Similarly, promoters responsive to nutrients such as phosphate lack utility due to the requirement of plants for constant levels of these nutrients for normal growth in the

30 field.

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Several reports of attempts to regulate the expression of genes in transgenic plants have been reported. European patent application number 85302593.0 discloses the isolation of four heat shock gene promoters fr m soybean and claims their use for driving the expression of foreign genes in transgenic

	the application, the authors claim the
	use of these promoters in temporarily activating
5	expression of foreign genes such as a crystalline
	toxic protein structural gene of Bacillus
	thuringensis or an herbicide resistance gene in
	response to heat stress in vivo. However, this
	leaves the expression of a gene linked to one of
10	these heat shock promoters to chance changes of the
	daily temperature in the field.
	Marcotte and Quatrano [J. Cellular Biochem,
	Supplement 12C, 1988; Marcotte, W. R., Bayley, C. C.,
	and Quatrano, R. S., Nature 335, 454-457 (1988)] have
15	reported initial results of studies of the
	inducibility of a chimeric gene whose transcription
	is driven by promoter fragments derived from two
	abscisic acid (ABA)-inducible genes (Em and a 75
	globulin) from wheat. The products of these genes
20	were shown to be induced in whole plants by addition
	of ABA. The induction was shown to be, at least in
	part, at the level of transcription. Promoter
	fragments of varying lengths from the 5' region an Em
	genomic clone were translationally fused to a
25	bacterial G-gluquenida (gua)
	bacterial β-glucuronidase (GUS) coding region that
	was linked to polyadenylation signals from the CaMV
	358 transcript. The ABA inducibility of GUS activity
	using these different length promoter fragments was
30	analyzed in transient expression assays using both
	monocot (rice) and dicot (tobacco) protoplasts. They
	demonstrated that regions upstream of the Em coding
	region (650 bp) and the 7S globulin coding region
	(1800 bp) contain sequences that are sufficient for
	ABA-regulated expression of GUS activity in rice

35 protoplasts transient assays. The Em promoter failed to show any responsiveness in the dicot transient

expression assay, indicating that the promoter may
not function in dicot plant species. However, as
discussed in detail in an earlier section of this
work, the induction of undesirable pleiotropic
effects resulting from application of phytohormones
(including ABA) to whole plants in the field
precludes the use of these compounds in regulating
gene expression in transformed plants.
A patent was issued in Europe to De Danske
Sukkerfab A/B [CC87-106623] that claims a method to
improve the nitrogen fixing system of leguminous
plants by controlling the expression of genes of
interest with a promoter from a root/ nodule specific
gene. Specifically, the inventors demonstrated that
a chloramphenicol acetyltransferase (CAT) gene driven
by the promoter derived from a soybean leghemoglobin
gene was inducible in the roots of transformed plants
in a fashion similar to other root specific genes
that are affected by nodulation. The method is
severely limited in that induction of genes is
limited to simulation by nodulation and the induction
is root specific. It comes and the induction
is root specific. It cannot provide a true means to
externally control the expression of genes at any
time in all tissues of field grown transformed plants.
To date, there are no reports of practical
means to externally regulate the expression of
foreign genes in transgenic plants using a method
compatible with those used in normal agronomic
practices. While reports of plant promoter sequences
stimulated by light, heat, anaerobic stress, and
phytohormones have appeared, no disclosures of
specific inducible promoters that are responsive to
- mar are responsive to

chemical substances that might constitute the basis

for a practical method to control gene expression in

plants by application of the chemical in the field have appeared. At this time, a clear need exists for such promoter sequences to be used in recombinant DNA constructions that would enable one to externally control the expression of genes that can confer agronomic advantages if expressed at the proper time. Further, this specificity of expression should be amenable to external control through exposure of plants to chemical substances which can be readily.

plants to chemical substances which can be readily applied by a variety of application methods and which only induce the expression of the desired target gene.

SUMMARY OF THE INVENTION

A practical means to control the expression of selected genes in transformed plants and plant tissues by the application of a chemical substance has been discovered. The present invention provides nucleic acid promoter fragments and downstream sequences derived from corn, tobacco and petunia genes whose expression are responsive to a number of substituted benzenesulfonamides, and other compounds. These nucleic acid promoter fragments have been incorporated into recombinant DNA constructs containing a structural gene of non-plant origin. Transformation of plants with such constructions demonstrate that the expression level of the structural gene is regulated by chemical treatment. Specifically, one aspect of the present invention is a nucleic acid promoter fragment inducible by a compound of Formula I-IX:

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VII

VIII

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ΙX

wherein

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X is H, F, C1, Br CF₃, or C_1 - C_2 alky1; x^1 is H, F, C1, C_1 - C_2 alky1, $SO_2NR^1R^2$ or CO_2R^1 ; Y is H, C1 or $SO_2NR^1R^2$, CO_2R^1 , NO_2 , $P(O)(OR^1)_2$; R is H, C1-C6 alkyl, C3-C6 cycloalkyl, benzyl or C2-C4 haloalkyl or C2-C4 substituted with C1-C2 alkoxy or C1-C2 alkylthio;

R1 is C1-C3 alkyl;

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R^2 is C_1-C_3 alky1;
             R3 is CO2R2;
   5
             R^4 is C_1-C_6 alkyl or C_3-C_6 cycloalkyl;
             R5 is C1-C3 alkoxy or NR6R7;
             R^6 is H, OCH<sub>3</sub>, C_1-C_4 alkyl, C_3-C_6 cycloalkyl,
                C_1-C_4 alkyl substituted with C_1-C_2 alkoxy or
                ethoxyethoxy; and
 10
             R7 is H or C1-C2 alkyl;
      and agriculturally suitable salts thereof such that
      exposure of plants transformed with said promoter
      fragment to a compound of Formula I-IX causes
      increased expression of a DNA sequence coding for a
 15
      selected gene product operably linked to said
      promoter fragment.
            Preferred nucleic acid promoter fragments are
      obtained from plants, while more preferred nucleic
      acid promoter fragments are obtained from
 20
      monocotyledenous plants including corn, oats, millet,
      wheat, straw, barley, sorghum, amaranth, onion,
      asparagus and sugar cane; and from dicotyledonous
      plant selected from the group consisting of alfalfa,
      soybean, petunia, cotton, sugarbeet, sunflower,
25
     carrot, celery, cabbage, cucumber, pepper, canola,
     tomato, potato, lentil, flax, broccoli, tobacco,
     bean, lettuce, oilseed rape, cauliflower, spinach,
     brussel sprout, artichoke, pea, okra, squash, kale,
     collard greens, tea and coffee. Most preferred are
30
     nucleic acid promoter fragments obtained from corn,
     specifically those homologous to cDNA clones 2-1,
     2-2, and 5-2.
           Preferred compounds by virtue of activity or
     ease of synthesis are compounds of Formula I wherein:
35
           X is H or 2-Cl;
           Y is 3-C1 or SO2N(CH3)2;
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R is H, C_1 - C_6 alkyl or C_5 - C_6 cycloalkyl; and

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compounds of Formula II wherein:
             R is C1-C4 alkyl or C5-C6 cycloalkyl;
  5
             R_4 is C_1-C_4 alkyl; and
      compounds of Formula III wherein:
             Rc is OCH2 or NR6R7;
            R<sub>6</sub> is H or C<sub>1</sub>-C<sub>4</sub> alkyl; and
             R- is H.
      More preferred for use with recombinant DNA
 10
      constructions whose expression is regulated by a 2-1
      promoter are the compounds N-(aminocarbony1)-
      2-chlorobenzenesulfonamide, 2-chloro-N-(methylamino-
      carbony1)benzenesulfonamide, 1-(n-buty1)-3-methy1-
      sulfonylurea, 1-cyclohexyl-3-(methylsunfonyl)urea,
 15
      diethyl [[2-(butylaminocarbonyl)aminosulfonyl]-
      phenyl]]phosphonate, methyl 1-[(aminocarbonyl)-
      aminosulfonyl]benzoate, 2,3-dichloro-N-[(cyclo-
      pentylamino)carbonyl]benzenesulfonamide, and
      N-(aminocarbony1)-2,3-dichlorobenzenesulfonamide.
 20
      Most preferred is N-(aminocarbony1)-2-chloro-
      benzenesulfonamide.
            More preferred for use with recombinant DNA
      constructions whose expression is regulated by a 2-2
     promoter are the compounds diethyl [[2-(butylamino-
25
     carbony1)aminosulfony1]pheny1]phosphonate, N'-[2-(n-
     butylaminocarbony1)]-6-chloro-N,N-dimethy1-
     1,2-benzenedisulfonamide, N-isopropylcarbamoy1-
     benzenesulfonamide, 2-chloro-N-(methylaminocarbonyl)-
30
     benzenesulfonamide, 2,5-dichloroacetanilide, N-(amino-
     carbonyl)-2-chlorobenzenesulfonamide, and
     1-cyclohexy1-3-(methylsulfonylurea. Most preferred
     is diethy1 [[2-[(butylaminocarbony1)aminosulfony1]-
     phenyl]]phosphat .
35
           More preferred for use with recombinant DNA
     constructions whose expression is regulated by a 5-2
```

		promoter are the compounds 2-chloro-N (methylamino-
		carbonyl)benzenesulfonamide, 1-(n-butyl)-3-methyl-
	5	sulfonylurea, methyl 2-[(aminocarbonyl)aminosulfonyl]-
•		benzoate, N-isopropylcarbamoylbenzenesulfonamide,
•.		N-(aminocarbonyl)-2-chlorobenzenesulfonamide and
		N'-[2-(n-butylaminocarbony1)]-6-chloro-N,N-dimethyl-
		1,2-benzenedisulfonamide. Most preferred is
	10	2-chloro-N-(mothyle-i-
		2-chloro-N-(methylaminocarbonyl)benzenesulfonamide.
		Another aspect of this invention involves a
		nucleic acid promoter fragment comprising a
		nucleotide sequence from the 5' flanking promoter
	15	regions of genes substantially homologous to specific
	13	Cones, such that exposure of plants to a
		with said promoter fragment to a compound of Fermila
		1-IA Causes increased expression of DNA sequence
		couling for selected gene products operable links
		the 3 end to said promoter fragment professes
	20	genes are those from corn homologous to come along
		2-1, 2-2, 218 or 5-2; those from petupia homeless
		to come cione P6.1; and those from tobacco bemele-
		to CDNA Cione TZ.I. Most preferred as a puglei
		promoter fragment for the regulation of orpassion of
	25	DNA sequences for selected gene products upon
		exposure to a compound of Formula I-IX are those
		derived from the corn 2-2 gene.
		Another aspect of the instant invention
		involves a recombinant DNA construct, capable of
	30	transforming a plant, comprising a nucleic acid
		promoter fragment of the invention, a DNA sequence
		coding for a selected gene product operably linked to
		said promoter fragment, and a suitable 3' downstream
۲,		region such that exposure of said transformed plant
	35	to a c mpound of Formula I-IX causes increased
		expression of said DNA sequence for a selected gene
		Tot a selected gene

30

35

time.

product. Preferred DNA sequences for selected geneproducts are those encoding for β-glucuronidase. genes encoding herbicide resistance such as mutant 5 acetolactate synthase and 5-enolpyruvylskikimate-3-phosphate synthase, genes encoding insect resistance, genes encoding protease inhibitors, genes encoding Bacillus thuringiensis insecticidal endotoxins. genes encoding phytohormone biosynthetic enzymes. 10 genes encoding ethylene biosynthetic enzymes, genes encoding auxin biosynthetic enzymes, genes encoding cytokinin biosynthetic enzymes, genes encoding giberellin biosynthetic enzymes, genes encoding 15 chitinases, genes encoding biosynthetic enzymes for oil production, genes encoding restriction endonucleases, genes encoding starch biosynthesis and/or degradation enzymes, denes encoding male sterility/fertility phenotype and genes encoding 20 transposors and/or transposessors. Yet another aspect of the invention involves

plants transformed with a recombinant DNA construct of the invention such that exposure of said transgenic plant to a compound of Formula I-IX causes increased expression of a DNA sequence coding for a selected gene product operably linked 3' to said promoter fragment. The seeds of such transgenic plants are also envisioned as embodiments of the invention.

A final aspect of the invention involves a

method of Causing increased expression of a selected gene product in a plant comprising the steps of (a) transforming said plant with a recombinant DNA construct described above, (b) exposing the transgenic plant to a compound of Formula I-IX, and (c) causing said transgenic plant to increase expression of said selected gene product at a desired

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the major steps used in one

5 preferred embodiment of the invention.

Figure 2 shows the nucleotide sequence of the 2-1 gene promoter from the gene designated as 21.14.

Figure 3 depicts the creation of plasmids pJE481-1(Nco I) and pJE484-62(Xba I) from the 21.14

10 corn gene.

pJE516.

Figure 4 shows subcloning of the 2-2 gene designated 2-2 #4 and the nucleotide sequence of the promoter from the 2-2 #4 gene.

Figure 5 shows the nucleotide sequence of the 15 5-2 gene promoter from the gene designated as 52.411.

Figure 6 depicts the creation of plasmid pMC75.j5 from the 5-2 corn gene.

Figure 7 shows the nucleotide sequence of the 218 gene promoter.

20 Figure 8 shows the nucleotide sequence and transcription start site of the petunia P6 gene 1 promoter from the gene designated as P6.1.

Figure 9 depicts the creation of plasmid P614 and P654.

25 Figure 10 depicts the creation of plasmid T217.
Figure 11 depicts the creation of plasmid

Figure 12 depicts the creation of plasmid pHPH220.

30 Figure 13 depicts the creation of plasmids pTDS130 and pTDS133.

Figure 14 depicts the creation of plasmid pTDS134.

Figure 15 depicts the creation of plasmid 35 pTDS231.

Figure 16 shows the nucleotide sequence of the 21.14 gene promoter indicating the positions of deletions mad in the promoter.

	24	
	Figure 17 depicts the creation of plasmid	
	pMC715.83.	
5	Figure 18 depicts the creation of plasmid	
	pMC7113.	•
	Figure 19 depicts the creation of plasmids	•
	P655, P657 and P658.	٠.
	Figure 20 depicts the creation of plasmid P660.	
10	Figure 21 shows the nucleotide sequence of the	
	443 promoter.	
	Figure 22 shows the nucleotide sequence of the 463 promoter.	
15	Figure 23 shows the nucleotide sequence of the 478 promoter.	
13		
	Figure 24 shows the nucleotide sequence of the	
	420 promoter,	
	Figure 25 depicts the creation of plasmid P627.	
	Figure 26 shows the results of RNAse protection	
20	analysis that demonstrates N-(aminocarbonyl)-2-chloro	
	benzenesulfonamide induction of the P6.1 gene in	
	transgenic tobacco.	

DETAILED DESCRIPTION OF THE INVENTION

Figure 27 depicts the creation of plasmids

Figure 28 depicts the creation of plasmid

P656, P661, P662 and P663.

pJE518 and pJE519.

The present invention provides DNA promoter
of SDMA promoter that are useful in bringing the expression
of DNA sequence coding for selected gene products
under the control of externally applied chemicals in
transgenic plants. The promoter fragments described
in this invention are derived from genes of corn,
tobacco, and petunia that were found to be strongly
inducible by a number of substituted benzene-

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٠,

herbicide antidotes. Expression of the gene product is obtained by treatment of the transgenic plant with a suitable inducing compound. To accomplish the invention, cDNA libraries were made using RNA from the roots of plants treated hydroponically with the chemical N-(aminocarbonyl)-2-chlorobenzenesulfonamide, a compound of formula I wherein X is H, Y is Cl, and R is H. Libraries were differentially screened using a strategy designed to identify clones representing mRNA species whose steady-state levels rise following treatment with

this compound. These cDNAs were then characterized and used as hybridization probes to isolate the gene(s) encoding the induced RNAs from appropriate libraries of plant genomic DNA. Comparison of the nucleotide sequences derived for the cDNAs and their

20 corresponding genomic clones permitted identification of putative promoter, structural gene, end 3' downstream regions for each gene. The DNA fragments comprising the promoter regions from these genes were isolated and operably linked to foreign coding regions to create novel chemically inducible genes.

Suitable 3' downstream regions containing polyadenylation signals were added to the promoter/coding region fusions to complete the construction of chemically inducible recombinant

30 genes. These genes were then transformed into both plants and plant-derived tissues. Assays of N-(aminocarbonyl)-2-chlorobenzenesulfonamide-treated plants and plant tissues transformed with these DNA constructions dem

15 functional in transgenic plants and that they retain their responsiveness to ext rnal chemical stimulation.

5	have been found to have promoters that are inducible
5	by a number of compounds of formulae I-IX, it is
	likely that any number of plant species will possess
	promoters responsive to selected members of these
	classes of chemistry. Therefore, it is expected that
	the invention can also be accomplished using
10	promoters unrelated to those disclosed here that are
	derived from other plant species as long as the
	expression of the promoter is responsive to scope of
	chemistry defined in this invention. Indeed, it is
	expected that the invention may well be accomplished
15	by using promoters derived from genes inducible by
	compounds of formulae I-IX that are isolated from any
	prokaryotic or eukaryotic species.
	The promoters disclosed in this work may be
	further modified if desired to alter their expression
20	characteristics. It is expected that a small DNA
	fragment can be derived from a chemically-inducible
	promoter that is responsible for the chemical
	responsiveness of that promoter. This fragment may
	be combined with suitable regions from other
25	promoters to create recombinant promoters whose
	expression level can be increased in transformed
	plants by treatment with compounds of Formulae I-IX.
	For example, the 77 bp fragment corresponding to
	bases 264 and 340 of Figure 4 that appears to be
30	necessary for chemical responsiveness in the 2-2
	promoter may be incorporated into seed-specific
	promoters such as the β-conglycinin or phaseolin
	promoters to create chimeric promoters that are
	chemically inducible and active only in developing

35 seeds. Similarly, any number of chimeric promoters can be created by ligating a DNA fragment sufficient

		to confer chemical inducibility from any of the
		promoter claimed here to constitute promoters or
	5	promotes with other specificities such as
•		tissue-specific promoters, developmentally-regulated
4,		promoters, light-regulated promoters, stress-
		responsive promoters, hormone-responsive promoters
		and so on. This should result in the creation of
	10	chimeric promoters capable of inducing expression of
		gene products in any plant tissues or combination of
		tissues at any specific time in the plant's life
		cycle in response to chemical treatment.
		Chemically-inducible promoters disclosed herein
	15	include possible variations of said promoters such as
		those derived from deletion, rearrangement, random or
		controlled mutagenesis of the promoters, promoters
		driven by ligation with foreign operator regions,
		promoters ligated to enhancer or enhancer-like
	20	elements (transcription activators) from any source
		such as the enhancer-like element from the 35S
		cauliflower mosaic virus transcripts, etc.
		It is believed that any 3' downstream region
		capable of providing a polyadenylation signal and
	25	other regulatory sequences that may be required for
		the proper expression and processing of a mRNA may be
		operably linked to the 3' end of a structural gene to
		accomplish the invention. This would include the
		native 3' end of the homologous gene from which the
	30	chemically-inducible promoter itself was derived
		the 3' end from a heterologous gene encoding the same
		protein in another species, the 3' end from viral
•		genes such as the 3' end of the 35S or the 19S
•		cauliflow r mosaic virus transcripts, the 3' end of
•	35	the opine synthesis genes of Agrobacterium
		tumefaciens, the 3' ends of RUBISCO or CAB genes, or

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PCT/US90/01210 28 the 3' end sequences from any source such that the sequence employed provides the necessary regulatory information within its nucleic acid sequence to result in the proper expression of the promoter/coding region combination to which it is operably linked. Since the transcription start site for each of the various genes diclosed in this work has yet to be determined for all promoters the numbers for nucleotide positions in the various promoter fragments used in constructions are based upon either the assignment of the A residue of the ATG codon that initiates translation of the protein encoded by that gene as nucleotide 1 of the promoter fragment or assignment of the actual transcription start site as nucleotide 1. Nucleotides 5' to number 1 residue are numbered sequentially starting with -1. It is understood and expected that the DNA sequence between the transcription start site in each of these promoter fragments and the translation start site.

i.e. the region comprising the 5' untranslated leaders of the mRNAs encoded by these genes, can be replaced by other 5' untranslated leaders from other genes without affecting the chemical-inducibility of the resulting DNA constructions. In the context of this disclosure, a number of

terms shall be utilized. As used herein, the terms "promoter" and "promoter region" refer to a sequence of DNA, usually upstream (5') to the coding sequence of a structural gene, which controls the expression of the coding region by providing the recognition for RNA polymerase and/ r other factors required for transcription to start at the correct site. sequences are necessary but not always sufficient to

	drive the expression of the gene. A "promoter
	fragment" constitutes a fraction of the DNA sequence
5	of the promoter region. "Nucleic acid" refers to a
	large molecule which can be single stranded or double
•	stranded, composed of monomers (nucleotides)
•	containing a sugar, phosphate and either a purine or
	pyrimidine. In higher plants, deoxyribonucleic acid
10	(DNA) is the genetic metalic, deoxyribonucleic acid
	(DNA) is the genetic material while ribonucleic acid (RNA) is involved in the translation of the
	information from DNA into proteins. The term
	"nucleotide sequence" refers to a polymer of DNA or
	RNA which can be single- or double-stranded,
15	ortionally containing synthetic, non-natural or
	altered nucleotide bases capable of incorporation
	into DNA or RNA polymers. As used herein, "DNA
	sequence for a selected gene product" refers to a DNA
	sequence that codes for a specific RNA transcript.
20	"Suitable regulatory sequence", as used herein,
	refers to a nucleotide sequence located upstream
	(5'), within, and/or downstream (3') to a DNA
	sequence for a selected gene product whose
	transcription and expression is controlled by the
25	regulatory sequence, potentially in conjunction with
	the protein biosynthetic apparatus of the cell. "RNA
	transcript" refers to the product resulting from the
	RNA polymerase catalyzed transcription of a DNA
	sequence. The RNA transcript may be a perfect
30	complementary copy of the DNA sequence and is
	referred to an the minute of sequence and is
	referred to as the primary transcript or it may be an
4	RNA sequence derived from posttranscriptional
•	processing of the primary transcript and is referred
1	to as the mature RNA. "Regulation" and "regulate"
, 35	refer to the modulation of gene expression induced by

DNA sequence elements located primarily, but not

	exclusively upstream of (5' to) the transcription	
_	start of a gene. Regulation may regulation and	
5	none response to a stimulation, or it many	,
	variations in the level of gene expression	
	"Responsive" and "response", as used herein, refer to	
	the change in the expression level of a regulated	٠,
	promoter or gene following the application of an	
10	environmental stimulus. The term "structural" gene	
	refers to that postions. The term "structural" gene	
	refers to that portion of a gene encoding a protein,	
	polypeptide, or a portion thereof, and excluding the	
	regulatory sequences which drive the initiation of	
15	transcription. A structural gene may be one normally	
	Tound in the Cell Of it may be one met	
	d Cellular location wherein it is interest.	
	which case it is termed a heterologous ser-	
	Meterologous gene may be derived in whole and	
	any source known to the art, including	
20	pacterial genome or episome, eukarvotic puel-	
	presente DNA, CDNA, Or Chemically synthogical and	
	The structural gene may constitute an uninterrupted	
	coding region or it may include one or more introns	
25	structural gene may be a composite of segments	
	derived from different	
	derived from different sources, naturally occurring	
	or synthetic. A "3' downstream region" (or "3'	
	end") refers to that portion of a gene comprising a	
30	our segment, excluding the 5' seguence which a	
30	the initiation of transcription and the atomic	
	portion of the gene, that contain a polyadamai.	
	signal and any other regulatory signals	•
	affecting mana processing or gene expression my	•
	polyadenylation signal is usually characteries?	
35	attecting the addition of polyadenylic said	•
	the 3' end of the mRNA precursor. Polyadenylation	
	- Lolyadenylation	

signals are commonly recognized by the presence of homology to the canonical form 5'-AATAAA-3', although variations are not uncommon. The term "recombinant DNA construct" refers to a plasmid, virus, autonomously replication sequence, phage or nucleotide sequence, linear or circular, of a singleor double-stranded DNA or RNA, derived from any 10 source, in which a number of nucleotide sequences have been joined or recombined into a unique construction which is capable of introducing a promoter fragment and DNA sequence for a selected gene product along with appropriate 3' untranslated 15 sequence into a plant cell. As used herein, "plant" refers to whole plants and plant-derived tissues. "Plant-derived tissues" refers to differentiated and undifferentiated tissues of plants, including, but not limited to roots, shoots, leaves, pollen, ovules, 20 seeds, tumor tissue, and various forms of cells in culture such as intact cells, protoplasts, embryos and callus tissue. Plant-derived tissues may be in planta or in organ, tissue or cell culture. A "monocotyledonous plant" refers to a plant whose 25 seeds have only one cotyledon, or organ of the embryo that stores and absorbs food. A "dicotyledonous plant" refers to a plant whose seeds have two cotyledons. A "protoplast" refers to a plant cell without a cell wall or extracellular matrix. As used 30 herein, "transformation" means processes by which cell/tissue/plant acquire properties encoded on a nucleic acid molecule that has been transferred to the cell/tissue/plant. "Transferring" refers to methods to transfer DNA into cells including 35 microinjection, or permeabilizing the cell membrane with various physical (e.g., electroporation) or

	chemical (e.g., polyethylene glycol, PEG) treat-
	ments. As used herein, "exposure of" a protoplast or
5	a plant to a chemical substance refers to treating,
	incubating, contacting said protoplast or plant with
	the substance. The term, "operably linked" refers to
	the chemical fusion of two fragments of DNA in a
	proper orientation and reading frame to be trans-
10	cribed into functional RNA. As used herein, the term
	"homologous to" refers to the similarity between the
	nucleotide sequences of two nucleic acid molecules or
	between the amino acid sequences of two protein mole-
	cules. Estimates of such homology are provided by
15	the use of either DNA-DNA or DNA-RNA hybridization
	under conditions of stringency as is well understood
	to those skilled in the art (as described in Hames
	and Higgins (eds.) Nucleic Acid Hybridization, IKL
	Press, Oxford, UK]; or by the comparison of the se-
20	quence similarity between two nucleic acids or pro-
	teins. As used herein, "substantially homologous"
	refers to nucleic acid molecules which require less
	stringent conditions for hybridization than condi-
	tions required for such molecules to be homologous to
25	each other; as well as to DNA protein coding se-
	quences which may involve base changes that do not
	cause a change in the encoded amino acid, or which
	involve base changes which may alter an amino acid
	but not affect the functional properties of the pro-
30	tein encoded by the DNA sequence, or this may refer
	to DNA sequences involved in regulating transcription
	of a gene. Thus, the nucleic acid promoter fragments
	described herein include molecules which comprise
	possible variations of the nucleotide bases derived
35	from deletion, rearrangement, and random or
	controlled mutagenesis of the promoter fragment so
	long as the DNA sequences of the promoter fragments
	are substantially hom logous. "Effective sequence"

of a DNA sequence coding, for a protein refers to a
truncated version of the DNA sequence which encodes
peptide which is at least partially functional with
respect to the utility of the original protein. The
term "expression" as used herein is intended to mean
the transcription and/or translation to gene product
from a gene coding for the sequence of the gene

- 10 product. In the expression, a DNA chain coding for the sequence of gene product is first transcribed to a complementary RNA which is often a messenger RNA and, then, the thus transcribed messenger RNA is translated into the above-mentioned gene product if the gene product is
- 15 the gene product is a protein. Expression, which is constitutive and further enhanced by an externally controlled promoter fragment thereby producing multiple copies of messenger RNA and large quantities of the selected gene product, is referred to as
- 20 "over-production". The "translation start codon" refers to a unit of three nucleotides (codon) in a nucleic acid that specifies the initiation protein synthesis.
- The techniques of DNA recombination used

 throughout this invention are known to those skilled in the art and are generally described in Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, N. Y., 1982).

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Enzymatic Treatments of DNA

Restriction Enzyme Digestions

The restriction enzyme digestion buffers and digestion conditions used were those supplied by the manufacturer of each particular enzyme. Enzyme was added to give 5-10 units per micr gram of DNA and the

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reaction mixture was adjusted to the appropriate final volume with water (usually 10-20 µ1). The restriction enzyme reaction mixtures used routinely contained 0.7-10 µg of plasmid DNA. The reaction mixtures were mixed and then generally incubated at the appropriate temperature for up to 2 hours. Digestion of DNA with multiple enzymes was done concomitantly when the optimal salt and temperature conditions of the separate enzymes are compatible. When these conditions were sufficiently different, digestions were done sequentially beginning with the enzyme requiring the lowest salt concentration. Subsequent reactions were supplemented to the

Gel Electrophoresis of DNA

For polyacrylamide gel electrophoresis of DNA,
20 the Tris-Borate-EDTA (TBE) buffer described by
Bethesda Research Laboratories, Gaithersburg, MD
20877 which consists of 89 mM Tris and 89 mM borate
(pH 8.3), 2.5 mM Na_EDTA was used. The gels used
consisted of 5% acrylamide and 0.2% bis-acrylamide
dissolved in 100 ml 1X TBE. To this solution, 0.225
ml of an aqueous 25% ammonium persulfate solution was
added.

appropriate buffer conditions for the enzyme used.

After adding 55 µl of N,N,N'.N'-tetramethyl ethylenediamine (TEMED), the solution was pipetted into a gel mold. One mm comb and spacers were commonly used and approximately 0.5 to 2 µg of DNA was loaded per well. Electrophoresis was carried out at 150 - 250 volts in 1 x TBE. After electrophoresis, the gel was stained in an agueous

35 solution of ethidium bromide (1 µg/ml) and the DNA was visualized on an ultraviolet transilluminat r. The gel was photographed using a Polaroid camera and Polaroid 667 film (Polaroid Tech. Photo, Cambridge,

5 MA 02139).

DNA was recovered from polyacrylamide gels as follows: The desired band, visualized by ethidium bromide (EtBr) staining, was cut from the gel, placed in an Eppendorf tube and minced with a teflon

- 10 pestle. An equal volume of a 0.5 M ammonium acetate, l mM EDTA solution was added and the tube was incubated at 37°C overnight with vigorous shaking. The following day, the tube was centrifuged at 14,000 x g in a microfuge for 10 minutes at room
- 15 temperature, the supernatant was removed, 1/2 volume of elution buffer was added to the minced polyacrylamide and the contents were mixed and vortexed. The tube was centricuged again as above.
- and the supernatant was removed and pooled with the original sample. The pooled supernatants were passed over a small glass wool column to remove any residual polyacrylamide gel pieces and the DNA in the sample was precipitated by addition of 2 volumes of ethanol and incubation in dry ice-ethanol. The DNA was
- 25 collected by centrifugation of the sample in a microfuge, as above, for 15 minutes at 4°C. The pellet was then rinsed with 70% ethanol, dried under vacuum and resuspended in the buffer of choice depending on the nature of the next manipulation.

Agarose gel electrophoresis of DNA was performed in 0.7% agarose gels using the buffer described above for polyacrylamide gels.
Electrophoresis was conducted at a voltage of 50 to 150 volts depending on the amount of DNA per lane and the desired timing of the run. After electrophoresis, the gel was stained with 1 µg/ml of

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EtBr and the DNA is visualized on an ultraviolet transilluminator and photographed as described above. DNA was often recovered from agarose gels using low gelling temperature agarose, Sea Plaque Agarose from FMC Corporation, Marine Colloids Division, Rockland, ME 04841. The electrophoresis procedure was stated above. After visualization of the DNA of interest, the band was cut out and placed into a microcentrifuge tube. The tube was then frozen at -80°C for 30 minutes and then thawed. The agarose was then smashed with a pestle and the sample was centrifuged in a Beckman TL-100 table-top ultracentrifuge at 25,000 rpm for 30 minutes. The supernatant was removed from the tube without disturbing the agarose pellet at the bottom of the

tube. The sample was precipitated with the addition of 1/10 volume of 3 M sodium acetate pH 6.0 and 2 volumes of ethanol followed by a 15-30 minute incubation at -80°C. The DNA was recovered by centrifugation in a microfuge for 15 minutes at 4°C. The DNA pellet was then washed with 70% ethanol, dried under vacuum and resuspended in TE buffer.

Plasmid Isolation and Purification

A 25 ml overnight culture (or exponentially growing culture) of the bacteria containing the desired plasmid was prepared. Two ml of the overnight culture was diluted into 1 liter of M9CA or L broth (as described in Molecular Cloning: A Laboratory Manual, Maniatis T. et al., Cold Spring Harbor Laboratory, Cold Spring Harbor, NY) and incubated for 16 hours [overnight] at 37°C with vigorous shaking using appropriate antibiotic selecti n. The bacteria were collected by

	centrifugation at 4000xg [5500 rpm] in a GSA rotor]
	for 5 min at 4°C. The pellets were drained well and
5	resuspend in a total volume of 36 ml of GTE buffer
	(50 mM glucose, 25 mM TRIS-HCl, pH 8 and 10 mM
	The man gracese, 25 mar ikis-nci, ph 8 and 10 mM
	EDTA). Four ml of 40 mg/ml lysozyme were added to
	the bacterial suspension and the mixture was
	incubated at room temperature for ten minutes. The
10	cell suspension was cooled on ice and 80 ml of
	freshly made [0.2 N NaOH and 1% SDS] were added with
	gentle swirling to lyse bacteria. The lysate was
	incubated the on ice for 10 minutes 40 ml of 3 M
	potaggium agglete in 2 M agglet 40 ml of 3 M
15	potassium acetate in 2 M acetic acid were added. The
13	mixture was then incubated on ice for 15 minutes.
	The precipitate was removed by centrifugation at
	24,000g [12 K rpm] for 15 minutes and the supernatant
	was filtered through 4-5 layers of cheesecloth.
	Nucleic acids were precipitated by addition of 0.6
20	volumes of isopropanol. The resulting precipitate
	was collected by contribution of the resulting precipitate
	was collected by centrifugation at 12,000 rpm for 10
	minutes at 15°C in a GSA rotor. The pellet was
	washed with 70% ethanol (in TE buffer) and the DNA
	was re-centrifuged as before. The nucleic acid
25	pellet was dissolved in 3.85 ml of TE, pH 8. After
	the DNA has dissolved, 4.4 g of CsCl were added to
	the solution. After dissolution of CsCl, 0.32 ml of
	ethidium bromide (EtBr) was added to the solution
	from a 10 mg/ml stock (final concentration of 600
30	ug/ml). The plasmid DNA was banded by centrifugation
	at 65 000 rpm for at least 15 has
	at 65,000 rpm for at least 15 hr in a Beckman 70.1 Ti

while the two upper bands did absorb the dye. The 35 less dense top band which c rresponds to chromosomal DNA often was barely visible. The plasmid band,

rotor. The gradient generally contained three bands. The lowest band absorbed no ethidium bromide,

	which was the lower of the two EtBr absorbing bands
	was removed from the gradient by puncturing the side
5	of the tube below the band with a 20 gauge needle and
	drawing the DNA out of the tube. The EtBr was
	removed by repeated extraction of the DNA with NaCl
	saturated 2-propanol. This was made by adding 10 ml
	of 50 mM TRIS-HCl, pH 8.0, 1 mM EDTA and 10 ml of 5 M
10	NaCl to 80 ml of 2-propanol. The extracted plasmid
	DNA was diluted 3 fold with TE pH 8.0 and

precipitated with 2 volumes of ethanol at -20°C. The DNA was recovered by centrifugation at 10,000 g for 30 minutes, resuspend in TE buffer and

15 re-precipitated with sodium acetate and ethanol. The DNA was resuspend in TE buffer and stored at -20°C.

Biological Material Deposits

The following cell lines and plasmids, as

described herein, have been deposited with the

American Type Culture Collection, 12301 Parklawn

Drive, Rockville, Maryland 20852, and have been given

the following ATCC accession designations:

25			
23	ITEM	DATE	ATCC ACCESSION
	plasmid pln2-2-3 in E. coli strain HB101	9/27/88	67803
30	plasmid pIn5-2.32 in E. <u>coli</u> strain HB101	9/27/88	67804
	plasmid pIn2-1.12A in E. <u>Coli</u> strain HB101	9/27/88	67805
	plasmid pMSP ^T K in E. <u>coli</u> strain HB101	6/08/88	67723
35	plasmid T2.1 in	10/11/88	67822

plasmid P6.1 in E. coli JM83

10/11/88 67823

5 plasmid pJJ3431 in E. coli JM83

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2/03/89 67884

The present invention is further defined in the following EXAMPLES, in which all parts and percentages are by weight and degrees are Celsius, unless otherwise stated. It should be understood that these EXAMPLES, while indicating preferred embodiments of the invention, are given by way of illustration only. From the above discussion and these EXAMPLES, one skilled in the art can ascertain the essential characteristics of this invention, and without departing from the spirit and scope thereof, can make various changes and modifications of the invention to adapt it to various usages and

conditions. Further, the present invention is not to be limited in scope by the biological materials deposited, since the deposited materials are intended to provide illustrations of starting materials from which many embodiments of the invention may be derived. All such modifications are intended to fall

25 within the scope of the appended claims.

EXAMPLE 1

Identification, Isolation and Modification of the Promoter and 3' Downstream Regions of the 21,14 Corn 2-1 Gene

Growth and Chemical Treatment of Plants

Missouri 17 corn seeds were surface sterilized by soaking them in a solution of 10% commercial bleach and 0.1% sodium dodecylsulfate (SDS) for 30 minutes. Seeds were then rinsed thoroughly in a

	buchher runnel with sterile distilled water and
	prepared for germination by placing them onto 5-6
5	layers of moist sterile paper towels in a 8" x 10"
	glass baking tray. The tray was covered with
	aluminum foil and placed in the dark in a 30°C
	incubator for 48-72 hours to allow the seeds to
	germinate. After germination, seedlings were grown
10	hydroponically in an apparatus consisting of a sheet
	of 8 mesh stainless steel wire gauze suspended over
	the top of a 2 liter glass beaker filled with sterile
	half strength Hoagland's solution (referred to 0.5x
	Hoagland's) so that the roots extended through the
15	mesh and into the media. The hydroponic apparatus
	was aerated by introducing humidified air into the
	bottom of the beaker with a gas diffusing stone
	commonly used in tropical fish aquariums. The
	apparatus was covered with a loose-fitting sheet of
20	aluminum foil and placed in a reach-in growth chamber
	illuminated by both fluorescent and incandescent
	lamps at an intensity of 4400 lux. Seedlings were
	grown at 28°C, 75% relative humidity using a 16 hour
	day/8 hour night cycle. After two days, the foil was
25	removed and plants were grown for an additional 5-6
	days. Any 0.5% Hoagland's lost to evaporation was
	replenished every 2-3 days. On the tenth day, plants
	were transferred into either fresh 0.5% Hoagland's
	for untreated plants, 0.5% Hoagland's containing 0.2
30	g/liter of 2-chlorobenzenesulfonamide for chemically
	treated control plants, or 0.5% Hoagland's containing
	0.2 g/liter of N-(aminocarbony1)-2-chlorobenzene-
	sulfonamide for chemically treated plants. Plants
	were then allowed to grow for six additional hours
35	prior to harvest.

		Roots were harvested from hydroponically grown
	5	plants by removing the wire mesh from the beakers
		with the corn plants still intact. The roots were
ų		cut from the plants just below where they were
*		immersed in growth media and 10-15 g portions of root
	10	nicrogen. Frozen tissue was bear a
		"Z to a -out freezer where it was
		up to one year before use.
		Isolation of Total Cellular RNA From Root Tissue
		Submidine thiocyanate reagont
	15	the contents of a loo - L-LL
		Justice thiocyanate (Kodak Laborate
		10.6 ml 1 M Tris-HC1, pH 7.6 and 10.6 ml 200 mM
		Z Pil /. D. The solution was attached
	20	contents of the bottle were dissolved and 4.24 g of
		sodium lauryl sarcosinate and 2.1 ml
		β-mercaptoethanol were added. The volume of the
		solution was adjusted to any
		solution was adjusted to 212 ml with sterile H ₂ O and
	25	
		sterile filtration units. The guanidine thiocyanate
		and acored at 4°C in the dark until
		Frozen root tissue samples were removed from
		o cited and transferred to 1
	30	nitrogen. Once cooled to liquid N2 temperature,
		or tissue was transferred to a manta
		pestie that had been pre-cooled with 1:
,		and the cissue was ground to a fine mendant
		Fordered tissue was then transferred to
`.		corex centrifuge bottle containing since
		trans of fee cold guanidine thiographe
		ml of CHCl ₃ , 0.2 ml n-octanol, 1 drop pourite
		- utop pourite

(American Scientific Products, McGaw Park, IL 60085. CAT # B 1162-1), and 2.5 ml vanadyl ribonucleoside complex (Bethesda Research Laboratories. Gaithersburg, MD 20877, CAT # 5522SA). The tissue was then ground further by vigorous homogenization with a PT-10/35 polytron (Brinkmann Instruments) for one minute at maximum speed. The crude tissue 10 extract was then centrifuged at 27.000g for 10 minutes at 4°C. The supernatant was decanted into a graduated cylinder and 1 g of CsCl was added for each 2.5 ml of supernatant. The solution was then centrifuged at 36.000g for 10 minutes at 4°C and the 15 resulting supernatant was layered over 2 ml pads of 5.7 M CsCl (in 100 mM EDTA pH 7.6) in 9/16" x 3-1/2" polyallomar ultracentrifuge tubes. The resulting step gradient was centrifuged at 35,000 rpm for 15 -20 hr at 10°C using a Beckman SW41Ti rotor or 20 equivalent. Following ultracentrifugation, the supernatant was carefully removed by aspiration and the tubes were inverted and allowed to drain well. With the tubes still inverted, the tops of the tubes were cut off using a razor blade and discarded, 25 saving only the bottom 1.5 cm containing the RNA pellets. The sides were carefully wiped clean with a labroatory tissue wipe and the pellets were dissolved in 0.2 ml of TES buffer (10 mM TRIS-C1 pH 7.4, 5 mM EDTA, 1% SDS) and transferred to a 15 ml Corex® 30 centrifuge tube. The bottom of each polyallomar tube was rinsed with a second 0.2 ml aliquot of TES and then the two aliquots were combined. The RNA was

combined with an equal volume of chloroform:n-butanol (4:1 v/v) and vortexed briefly. The r sulting

emulsion was centrifuged at 8,000g for 5 min. at $20^{\circ}\mathrm{C}$ or at high speed in a clinical table-top centrifuge

for 10 minutes. The aqueous layer was transferred to a fresh 15 ml Corex® centrifuge tube, the organic

- phase was back-extracted with an equal volume of TES. and the two aqueous layers were pooled. RNA was precipitated at -20°C for at least 2 hr after adding a tenth volume of 3.0 M sodium acetate pH 6.0 and two volumes of ethanol. The RNA was recovered by
- 10 centrifugation at 10,000g for 20 min. at 4°C. The supernatant layer was cently aspirated off and the RNA was dissolved in 0.5 ml of either sterile water or 1 mM EDTA, pH 7.6. A small aliquot was diluted 100 fold with water and the A_{260} of this dilution 15 was measured to determine RNA concentration.

Isolation of poly(A) + RNA

Poly (A)+ RNA was purified from 5 mg of total cellular RNA preparations by thermoelution from

- poly-U-Sephadex®. All buffers were sterilized by 20 autoclaving prior to use. Total RNA was diluted to less than 500 $\mu g/ml$ with low salt poly-U buffer (20 mM Tris-Cl, pH 8.0, 1 mM EDTA and 0.1% SDS). The RNA was denatured by heating at 65°C for 5 minutes 25
- followed by rapid cooling on ice for 5 minutes. NaCl was added to a final concentration of 150 mM, and this solution was loaded onto a water jacketed column (Bio-Rad, 1414 Harbour Way South, Richmond, CA 94804, CAT # 737-2231) containing 2 g of poly U-Sephadex
- (Bethesda Research Laboratories, CAT # 5941SB) that had been equilibrated with high salt poly U buffer (20 mM Tris-Cl, pH 8.0, 1 mM EDTA, 0.1% SDS and 150 mM NaCl). The column was maintained at a temperature of 25-30°C with a circulating water bath. The column
- was then washed once with 6-7 ml of high salt poly-U 35 buffer. The running temperature of the column was

increased to 40°C and it was washed again with 6-7 ml of high salt poly-U buffer. Seven ml of low salt

- 5 poly-U buffer was then added to the column and the temperature was raised to 60°C. After waiting 5 minutes for the temperature of both the column and low salt poly-U buffer to equilibrate, poly (A)* RNA was eluted and collected in 0.5 ml fractions.
- 10 Fractions containing RNA (determined by measuring the A₂₆₀ of a small aliquot from each fraction) were pooled and ethanol precipitated as described earlier. RNA was re-precipitated as above but with potassium acetate rather than sodium acetate.
- 15 resuspended in water at a concentration of 1 mg/ml and stored at -80°C.

Construction of CDNA Libraries

cDNA was synthesized from 5 µg of

- N-(aminocarbony1)-2-chlorobenzenesulfonamide treated poly (A)* RNA using a cDNA synthesis kit (Amersham Corporation, CAT # RPN 1256). The manufacturer's recommended protocol was followed without modification. The mass of double-strended (ds) cDNA
- 25 synthesized was calculated from the amount of [α32P] dCTP incorporated during the first and second strand synthesis reactions. The average size of the cDNA
- synthesized was then estimated from its mobility during electrophoresis in alkaline agarose gels. The average number of 5' ends/ug of cDNA was then
- calculated. The double-stranded cDNA was ethanol precipitated and was recovered by centrifugation for 10 minutes at 4°C. The DNA pellet was briefly dried under vacuum and dissolved in H₂O. 250 uCi of [³H]
- 35 dCTP in 50% ethanol was added to a 1.5 ml microfuge tube and dried in vacuo. One microgram of ds cDNA in

CDNA.

		Fr was crombieried into this tube,
		followed by 25 µl of 2X tailing buffer (2.5 mM
	5	β-mercaptoethanol, 100 mg/ml BSA, 3.5 mM MnCl ₂ and
•		135 mM potassium cacodylate, pH 7.0). Ten units of
		terminal deoxynucleotidyl transferase was added and
		the tube was incubated at 30°C for 21 minutes. The
		tailing reaction was stopped by addition of EDTA to a
	10	final concentration of 20 mM and the tube was placed
		on ice. The C-tailed reaction products were
		extracted once with an equal volume of
		phenol:chloroform (1:v/v) and purified by spun-column
		chromatography. Spun column chromatography was
	15	performed by plugging the bottom of a 1 ml disposable
		syringe with sterile glass wool and filling it with
		Sephadex® G-50 that was equilibrated in STE buffer
		(TE, pH 8.0 containing 100 mM NaCl). The syrings was
		inserted into a de-capped 1.5 ml microfuge tube
	20	placed in the bottom of 15 ml Corex® centrifuge
		tube. The column was centrifuged at 1600g for 4
		minutes in a bench top clinical centrifuge.
		Additional Sephadex® G-50 was added and the column
		was spun again. This process was repeated until a
	25	packed bed volume of 0.9 ml was obtained. Two rinses
		of the column were conducted with 0.1 ml of STE
		buffer and the syringe was centrifuged as above
		between each rinse. DNA samples were loaded onto the
		column in a volume of 0.1 ml in STE buffer and the
	30	column was centrifuged in a decapped microfuge tube
		as described above. The DNA was recovered by
		collecting the effluent in a microfuge tube and
•		storing it at -20°C. The average number of dC
		residues added per 3' end of cDNA was then calculated
•	35	from the % incorporation of the [3H] dCTP into the

	Equilimolar amounts C-tailed ds cDNA and
	dG-tailed pBR322 vector DNA (New England Nuclear
5	Research Products, 549 Albany St., Boston, MA 02118
	CAT #NEE-118) were mixed together in 0.1 M NaCl, 10
	mM Tris-HCl, pH 7.8, and 1 mM EDTA in volume of less
	than 10 µl. The DNA in the mixture was annealed by
	first heating it to 70°C for 10 minutes in a water
10	bath. The bath was then turned off and the mixture
	was allowed to slowly cool to room temperature. The
	mixture was then moved to a cold room and slow-cooled
	to 4°C. Small aliquots of annealed DNA were used to
	transform competent E. coli HB101. Competent cells
15	were prepared by diluting 0.1 ml of an overnight
	culture of HB101 grown in LB broth into 50 ml of the
	same media. This fresh culture was grown at 37°C
	with shaking until it reached an A650 of 0.2-0.5.
	Cells were then harvested by centrifugation at 2500g
20	for 5 minutes at 4°C, resuspended in 25 ml of 0.25 M
	CaCl ₂ and kept on ice for 20 minutes. Cells were
	recovered by centrifugation as above, resuspeneded in
	0.5 ml of 0.1 M CaCl2, stored on ice and used within
	24 hours. One hundred microliter aliquots of these
25	competent cells were placed into sterile 4 ml
	polyproplylene tubes, mixed with aliquots of the
	annealing reaction from above, and the transformation
	mixture was incubated on ice for 15 minutes. The
	cells were then heat shocked for 5 minutes in a 37°C
30	water bath without shaking. The cells were returned
	to the ice for 2 minutes before addition of 2 ml of
	LB medium. The cells were then grown for one hour
	at 37°C with shaking and aliquots of the
	transformation mixture were plated on LR plates which

35 contained 12.5 µg/ml tetracycline (tet). Plates were

then incubated at 37°C overnight.

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Isolation of 2-1 cDNA clones

Antibiotic resistant colonies resulting from
the transformation were picked and arrayed onto 150
mm LB agar plates containing 12.5 µg/ml tet.
Colonies were grown up and transferred to 140 mm
nitrocellulose filters by layering pre-wetted filters
(accomplished by layering dry filters on fresh LB
plates containing 12.5 µg/ml tet) onto each plate.
The transferred colonies were grown up as above and
these filters were referred to as the master filters
of the cDNA library.

Two replica nitrocellulose filters were made of each master filter. To accomplish this, nitrocellulose filters were first prewetted as above. Individual wetted filters were then laid on top of a master filter and the pair of filters were placed between several sheets of dry 3MM paper This

sendwich was placed between two glass plates that were then were pressed together to transfer bacteria from the master filter to the replica. The filters were then separated and the replica was place on a fresh LB/tet plate. This process was repeated until

25 two replicas of each master filter had been made Master filters were returned to fresh plates and stored at 4°C.

Replica filters were grown at 37°C until

colonies reached 1-2 mm in diameter and then filters were transferred onto LB plates containing 200 µg/ml chloramphenicol. The plates were incubated overnight at 37°C. The next morning, bacteria on the filters were lysed and their DNA was fixed to the filters in gitu. To lyse bacteria, filters were

35 removed from the agar plates and placed colony side up for three minutes in a glass tray containing 3

sheets of Whatman 3MM paper that had been saturated with 10 % SDS. Filters were then transferred for 5 minutes to a tray containing 3 sheets of Whatman 3MM paper saturated with 0.5 N NaOH, 1.5 M NaCl, followed by transfer to a tray containing 3MM paper saturated with 1 M Tris-HCl pH 7.5, 1.5 M NaCl for 6 minutes. The filters were air dried for one hour and baked at 10 70°C for 2 hours in vacuo. Replica copies of the cDNA library were differentially screened for clones representing mRNAs whose abundance rise following N-(amincarbonyl)-2-chlorobenzenesulfonamide treatment. To accomplish this, one replica of each master filter was 15 hybridized with a 32P-labeled single-stranded cDNA probe made by reverse transcribing poly(A)+ RNA from untrested corn roots, while the other replica filter was hybridized with a 32p single-stranded cDNA probe 20 made by reverse transcribing poly (A)+ RNA from N-(aminocarbonyl)-2-chlorobenzenesulfonamide-treated corn roots. Probes were synthesized from 5 µg of each poly(A)+ RNA by performing first strand cDNA synthesis using the Amersham cDNA synthesis kit. 25 First strand reactions were terminated by addition of EDTA to 20 mM, and then NaOH was added to a final concentration of 0.4 M to hydrolyze RNA. After RNA hydrolysis had been carried out for 6 hours at 22°C. the pH of the cDNA solution was adjusted to 30 neutrality with HCl and the first strand reactions were applied to a 1 cm X 10 cm Sephadex® G-100 column that was equilibrated with 10 mM Tris-HCl, pH 8.0, 20 mM NaCl, 0.2% SDS. Radiosstive material eluting in the void volume was pooled and the DNA was ethanol

precipitated. Labeled DNA was collected by centrifugation at 14,000g for 20 minutes at 4°C. The

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in a so	1011	volume	T	HG 3	8	0.	The	F 2 4 4		ivitu
pellet	was	dried	in	Yacı	1Q	and	the	DNA	was	resuspended

5 incorporated into the probe was determined by counting a 1 µl aliquot in a liquid scintillation counter using 5 ml of scintillation fluid.

Replice filters were divided into two sets of filters such that each set represented one copy of the containing the containing the containing the containing the colony sides facing outward. Each bag was filled with 70 ml of hybridization buffer, sealed, and incubated overnight at 65°C in a water bath. Hybridization

buffer consists of 6X SEC (IX SEC is 0.15 M NaCl, 0.015 M trisodium citrate, pH 7.0), 2X Denherdt's (Denhardt's is 0.02% bovine serum albumin (BSA), 0.02% polyvinyl pyrrolidine, 0.02% Ficoll Type 400

(MWr 400,000), 0.5 % SDS, 50 mM sodium phosphate pH 20 6.8, 2 mM EDTA and 100 μg/ml denatured calf thymus DNA.

Screening of the library was accomplished by discarding the hybridization buffer in each bag and replacing it with 30 ml of hybridization buffer

- 25 containing 5 x 10⁶ cpm/ml of probe made from poly(A)+
 RNA purified from corn root tissue which had been
 treated with N-(aminocarbony1)-2-chlorobenzenesulfonamide for six hours in the hydroponic
 system. The filters representing the second copy of
- 30 the library was hybridized in the same manner with 5 X 10⁶ cpm/ml of probe made from poly(A)⁴ RNA isolated from roots of plants that had not been treated. The filters were hybridized at 65°C for a minimum of 48 hours. Filters were then removed from the bags and
- 35 washed twice for 15 minutes at room temperature with 2X SSC, 1 mM EDTA, 0.2% SDS and 1 mM sodium

pyrophosphate, once at 65°C with a 0.5 X SSC and 0.1% SDS for one hour, and once for thirty minutes at 65°C 5 with 0.2 X SSC and 0.1% SDS. Filters were air-dried briefly and exposed to Kodak XAR-5 film at -80°C for approximately 36 hours using a single Du Pont Lightning Plus intensifying screen. Autoradiograms of the filters were developed using a Kodak automated 10 film processor. Any colony displaying a stronger hybridization signal with the probe made using RNA from N-(aminocarbonvl)-2-chlorobenzenesulfonamidetreated plants than with the probe made using RNA from untreated RNA was deemed a positive clone in the 15 differential screen and selected for further analysis. One colony from the differential screen. designated 2-1, was chosen as a potential positive

clone and was chosen for further analysis. Plasmid DNA was prepared from the 2-1 colony using a small 20 scale plasmid DNA isolation procedure. This was accomplished by inoculating 5 ml of LB medium containing the appropriate antibiotic (tet) with the single bacterial colony. After overnight incubation at 37°C with vigorous shaking, 1.5 ml of the culture 25 was poured into a microcentrifuge tube. The tube was centrifuged for 20 seconds in a microcentrifuge and the medium was removed by aspiration leaving the bacterial pellet as dry as possible. An additional 1.5 ml of culture was added to the tube and the above 30 steps were repeated. The pellet was resuspended in 100 µl of an ice-cold solution of GTE buffer (50 mM glucose, 10 mM EDTA, 25 mM TRIS-HCl, pH 8.0) with 4 mg/ml lysozyme (added to the solution just before use) with vortexing. After 5 minutes at room 35 temperature, 200 ml of a freshly prepared solution of

lysis buffer (0.2 N NaOH and 1% SDS) was added t the

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tube and the contents were mixed by inverting the tube rapidly two or three times. The tube was placed on ice for 5 minutes, followed by addition of 150 ul of an ice-cold solution of potassium acetate pH 4.8 (made by adding 11.5 ml of glacial acetic acid and ٠ 28.5 ml of $\rm H_2O$ to 60 ml of 5 M potassium acetate). The contents were mixed by inverting the tube sharply 10 several times. After 5 minutes on ice, the tube was centrifuged for 5 minutes in a microcentrifuge at 4°C. The supernatant was transferred to a fresh tube and an equal volume of phenol:chloroform (1:1 v/v) was added with mixing. The resulting emulsion was 15 centrifuged for 2 minutes in a microcentrifuge and the supernatant was transferred to a fresh tube. Two volumes of ethanol were added and the contents of the tube were mixed well. After 2 minutes at room temperature, DNA was collected by centrifugation for 20 5 minutes in a microcentrifuge. The supernatant was discarded and the tube was stood in an inverted position on a paper towel to allow all of the fluid to drain away. The pellet was washed with 250 μl of 70% ethanol and the tube was then recentrifuged. The 25 supernatant was discarded and the pellet was dried briefly in vacuo. Crude plasmid DNA was dissolved in 50 μ l of TE pH 8.0. The plasmid contained within clone In 2-1 was designated pIn 2-1.

An aliquot of the plasmid preparation was labelled by nick- translation using a commercial kit (Bethesda Research Laboratories, CAT# 81605B) following the manufacturer's suggested protocol. The labeled DNA was purified from the unincorporated nucleotides by spun column chromatagraphy.

An RNA slot blot procedure was used to confirm that the putative positive clon isolated during the

screening of the cDNA library represented an mRNA that was strongly induced by N-(aminocarbony1)-5 2-chlorobenzenesulfonamide. A nitrocellulose filter (Schlicher and Schull BA-85) was wetted by soaking it twice for 10 minutes in water, followed by a 10 minute soak in 1 M ammonium acetate. The filter was then placed into a Slot Blot apparatus (Schleicher and Schuell, Inc., Keene, NH 03431, CAT # 10 SRC072/0). Multiple 2.5 µg samples of total RNA from untreated corn roots, roots treated with 2-chlorobenzenesulfonamide, and roots treated with N-(aminocarbonyl)-2-chlorobenzenesulfonamide were 15 diluted to a final volume of 80 µl with sterile water. Forty ul of denaturation buffer (30% formaldehyde, 100 mM sodium phosphate pH 6.8) were added to each sample and all samples were then incubated at 65°C for 20-30 minutes and quick-cooled 20 in an ice slurry for 5 minutes. Thirty μl of 4 M ammonium acetate were added to each sample and the 150 ul samples were added to slots in the blotting cell with the aid of a 10-15 mm Hg vacuum. The filter was removed from the blotting cell, air dried and baked for 2 hours at 70°C in vacuo. 25 The filter was cut into six pieces such that each piece had one slot containing RNA from each of the three treatments described above. One of the filter pieces was incubated with 10 ml of 30 prehybridization buffer (50% deionized formamide, 5X SSC, 5X Denhardt's, 100 µg/ml denatured calf thymus DNA, 20 µg/ml homopoly(A), 40 mM sodium phosphate pH 6.8 and 0.5% BSA) in a heat-sealable bag for 6 hours at 42°C with occasional mixing. The filter piece was

then hybridized with nick-translated pIn 2-1. This

was performed by discarding the prehybridization

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solution from the bag and replacing it with 2.5 ml of hybridization buffer (50% deionized formamide, 5X SSC, 100 μg/ml denatured calf thymus DNA, 20 μg/ml homopoly(A) and 40 mM sodium phosphate, pH 6.8) containing 1.25 X 107 cpm of nick translated 2-1 plasmid described above. Nick-translated plasmid was denatured by boiling for 10 minutes followed by 10 quick-cooling on ice. The filter was then hybridized overnight at 42°C with occasional mixing. The filter was removed from the bag and washed twice at room temperature for 10-15 minutes on a rocking shaker with 2X SSC, 1 mM EDTA, 20 mM sodium 15 phosphate pH 6.8, 1 mM sodium pyrophosphate and 0.5% SDS and twice for 30 minutes at 65°C with 0.1% SSC and 0.5% SDS. The filter was briefly air-dried, wrapped in polyethylene food wrap and subjected to autoradiography overnight using Kodak XAR-5 film and 20 a single Du Pont Lightning Plus intensifying screen. The plasmid designated pIn 2-1 strongly hybridized to root RNA from N-(aminocarbonyl-2-chlorobenzenesulfonamide-treated plants, and hydridized extremely weakly, if at all, to RNA from 25 both untreated plants and 2-chlorobenzenesulfonamide-treated plants. By these criteria, cDNA clone 2-1 was confirmed as representing an mRNA induced by N-(aminocarbonyl)-2-chlorobenzenesulfonamide. 30 Plasmid pIn 2-1 was used as a probe in a northern analysis to determine the size of its corresponding mRNA. Two and a half μg of $poly(A)^+$ RNA from both untreated and N-(aminocarbonyl)-2-chlorobenzenesulfonamide-treated corn roots and 2.5 µg of Brome mosaic virus RNA (used as RNA molecular 35 weight markers) wer each placed in separate 1.5 ml

microfuge tubes, evaporated to dryness and taken up in 8 µl of Northern sample buffer (25% dejonized

- 5 formamide, 3% formaldehyde, 5 mM Na₂EDTA and 20 mM sodium phosphate pH 6.8). The RNA was incubated 15-20 minutes at 65°C, quick-cooled on ice, and 1 μ1 of northern loading buffer (5 mM sodium phosphate, pH 6.8, 50% glycerol and 0.2% bromophenol blue) was
- added to each tube. RNA samples were then loaded into 10 mm X 1 mm slots of a 1.5% agarose gel prepared in 20 mM sodium phosphate pH 6.8, 3% formaldehyde, and the RNA was subjected to overnight electrophoresis at 36-48 volts at room temperature in
- 15 10 mM sodium phosphate, pH 6.8, 3 % formaldehyde.
 The lanes containing BMV molecular weight

markers were cut from the gel with a razor blade and the remainder of the gel was blotted to a nylon membrane in a chemical fume hood essentially as

- 20 described by Thomas, P. S., Proc. Natl. Acad. Sci. USA, 77:520-5205 (1980). The agarose gel was inverted on a glass plate covered with two sheets of Whatman 3MM paper that had been saturated with 20X SSC. The glass plate was place over the top of a
- 25 baking dish filled with 20X SSC such that the ends of the 3MM paper extended over the edge of the glass plate and into 20X SSC in the dish. A sheet of Zeta-Probe nylon membrane (Bio-Rad Laboratories) was cut 0.5 cm larger than the gel, prewet in water, then
- 30 soaked for several minutes in 20X SSC. The membrane was laid on top of the gel and covered with a sheet of Whatman 541 paper soaked in 20X SSC followed by and several sheets of 3MM paper soaked in 20X SSC. A 10 cm stack of paper towels was then placed on top of
- 35 the 3 MM sheets to draw buffer through the gel, and RNA in the gel was transferred to the membrane

overnight at room temperature. The resulting RNA

— blet was then removed from the top of the gel after
marking the positions of the sample wells of the gel
relative to the membrane. The filter was air-dried
for one hour and then baked for 2 hours at 70°C in

- relative to the membrane. The filter was air-dried for one hour and then baked for 2 hours at 70°C in <u>vacuo</u>.

 The RNA molecular weight markers were stained in 100 mM NaCl 1 un/ml of Park for the RNA RNACL 1 un/ml of Park for the RNACL 1 un/ml of Park for the
- 10 in 100 mM NaCl, l μg/ml of EtBr for 1-2 hours followed by destaining with shaking in 100 mM ammonium acetate, 10 mM β-mercaptoethanol for 2-3 hours. The positions of the RNA markers were recorded by photographing the gel on an ultraviolet transilluminator. The migration distances of each RNA molecular weight marker was plotted against the log of its molecular weight to establish a standard
- curve. This standard curve was used to estimate the size of the 2-1 mRNA by its position in the same 20 agarose gel.

The RNA blot was prehybridized in Northern prehybridization buffer (50% deionized formamide, 5% SSC, 5% Denhardt's, 100 µg/ml boiled and sonicated calf thymus DNA, 20 µg/ml homopoly A, 40 mW sodium

- 25 phosphate pH 6.8 and 0.5% BSA) using 200 µl of buffer per cm2 of blot in a heat-sealed bag.
 Prehybridization was carried out for 6 hours at 42°C with occasional mixing. The plasmid pln 2-1 was nick-translated using a nick-translation kit as
- 30 described above to a specific activity of 5.9 x 108 CPm/μg of DNA. Prehybridization buffer was discarded and replaced with hybridization buffer (5% deionized formamide, 5x SSC, 100 μg/ml denatured calf thymus DNA, 20 μg/ml homopoly(A) and 40 mM sodium phosphate,
- 35 pH.6.8) containing 2 x 10⁵ cpm/ml of denatured, nick-translated pIn 2-1, using 100 µl of buffer/cm² of filter.

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The blot was hybridized for 24 hours at 42°C
with occasional mixing then washed twice at room
temperature for 10-15 minutes on a rocker with 2X
SSC, 5 mM Na₂EDTA, 25 mM sodium phosphate, pH 6.8,
1.5 mM sodium pyrophosphate and 0.5% SDS. This was
followed by two washes for 30 minutes each with 0.1X
SSC and 0.5% SDS at 64°C. The filter was air-dried,
wrapped in polyethylene food wrap and exposed
overnight to Kodak XAR-5 film at -80°C using a
single Du Pont Lightning Plus intensifying screen.
The Northern blot results were consistent with
those obtained in the slot blot experiment. No
hybridization was seen with untreated corn root RNA,
while a single intense hybridization signal to an
850-900 nucleotide (nt) mRNA was seen with

N-(aminocarbony1)-2-Chlorobenzenesulfonamide-treated RNA.

The size of the pIn 2-1 cDNA insert was analyzed by digesting the plasmid to completion with Pst I and subjecting the digestion products to

agarose gel electrophoresis. The results showed that pIn 2-1 insert was a single 450 bp Pst I fragment.

The pIn 2-1 insert not a full length copy of the message since Northern analysis indicated a 2-1 mRNA size of 850-900 nt. However, the pIn 2-1 insert was sufficiently large to use it as a probe for genomic clone isolations. A full-length cDNA clone was still

30 needed to determine the boundaries of the structural and regulatory regions of the 2-1 gene(s).
A new cDNA library was made from RNA isolated

from N-(aminocarbonyl)-2-chlorobenzenesulfonamidetreated corn roots using a procedure designed to maximize the probability of obtaining full length CDNA clones. First strand synthesis was performed in a 100 µl reaction containing 50 µg/ml of poly(A)+

RNA, 50 mM Tris-HC1, pH 8.3 at 42°C, 45 mM KC1, 0.5 mM GATP, GGTP and GTTP, 0.2 mM GCTP, 5 mM DTT, 7.5

- 5 μg/ml oligo (dT)12-18, 400 units/ml placental ribonuclease inhibitor, 7.5 mM MgCl₂, 4 mM sodium pyrophosphate, 0.4 mc/ml [a³²p] dCTP and 560 U/ml reverse transcriptase. The reaction was incubated at room temperature for 5 minutes and then transferred to 42°C for 45 minutes. The single strand cDNA was extracted sequentially with equal volumes of phenol, phenol:chloroform (1:1 v/v) and chloroform followed by ethanol precipitation in the presence of ammonium acetate.
 - The second strand was synthesized from 1 µg of first strand cDNA in a reaction containing 20 mM Tris-HCl, pH 7.5, 5 mM MGl₂, 10 mM (NH₄)₂SO₄, 100 mM KCl, 50 mg/ml BSA, 50 µM dNTPs, 0.1 mCi/ml [a32P] dCTP, 230 U/ml DNA polymerase I and 8.5 U/ml RNase
- 20 H. The reaction mixture was incubated for one hour at 12°C and one hour at 20°C. The products of the second strand reaction were size fractionated on a 1.0 x 15 cm Bio-Gel® A-50m (Bio-Rad Laboratories) column equilibrated and eluted with 0.3 M sodium
- 25 acetate in TE, pH 8.0. Fractions eluted from the column were collected and small aliquots of every second fraction were analyzed for cDNA size distribution by electrophoresis in a 1.2 % alkaline agarose gel. ³²P end-labeled Hind III digestion
- fragments of pUC18, pBR322 and SV40 were run in the gel as size markers. After electrophoresis, the DNA was fixed in the gel by soaking it in 15% TCA for 10-15 minutes. Excess liquid was removed from the gel by blotting to a stack of stacking weighted paper towards a placed on the paper.
- 35 towels placed over the gel for 1-2 hours and the gel

	was then wrapped in polyethylene wrap and exposed to
_	x-ray film. Column fractions containing cDNA greater
5	than 500 bp in length were pooled, ethanol
	precipitated twice, and dissolved in 8.5 ul of water
	Approximately 1-1.5 µg of cDNA was methylated
	at internal EcoRI sites by incubating it in 25 mm
	Tris-HC1, pH 7.5, 1 mM EDTA, 2.5 mM DTT, 10 µM
10	S-adenosylmethionine with 20 U of Eco RI methylase
	per microgram of cDNA at 37°C for 30 minutes. The
	methylase was inactivated by heating at 65°C for 10
	minutes and the DNA was extracted with
	phenol:chloroform (1:1) and precipitated with ethanol.
15	Eco RI linkers were were ligated to the cDNA by
	incubation of 2 µg of ds cDNA with 7.5 µg of
	phosphorylated linkers in 66 mM Tris-HCl, pH 7.5, 5
	mM MgCl ₂ , 5 mM DTT, 1 mM ATP and 20 units of T4 DNA
	ligase (New England Biolabs, Inc., Beverly, MA 01915,
20	CAT # 202). The reaction was incubated overnight at
	15°C. The products of the linker ligation reaction
	were digested to completion with 500 units of Eco RI
	for 4 hours at 37°C. The Eco RI digestion mixture
	was applied to a 1 X 10 cm Bio-Gel® A 50m column and
25	eluted with 0.3 M sodium acetate in TE, pH 8.0 to
	separate the cDNA from excess linkers and size
	alkaline agarose gel electrophoresis as described
30	above and fractions containing cDNA greater than 600
	bp were pooled and ethanol precipitated. The cDNA
	was resuspended in 100 µl of TE pH 8.0. The mass of
	CDNA was estimated by counting an aliquot of the the
	cDNA using the known specific activity of ³² P dCTP
25	used in the cDNA sysnthesis reactions. Aliquots of
35	the cDNA were then ligated to Eco RI digested

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and dephosphorylated lambda kgt 11 arms (Stratagene, 11099 North Torrey Pines Rd., LaJolla, CA 92037, CAT #200211) using ligation conditions described above. The ligation products were packaged with Gigapack Plus extracts (Stratagene) following the manufacturer's recommended protocol. The titer of

the resulting phage library was determined using E. coli Y1090 as a host. 10

Screening of Agt 11 Library

A 1.5 ml aligout of an exponentially growing culture of E. coli Y1090 grown in NZC broth were diluted with 0.6 ml of SM buffer (0.01% gelatin, 50 mM Tris-HCl pH 7.5, 5.8 g/l NaCl, 2g/l MgSO₄) and 2.1 ml of 10 mM MgCl $_2$, 10 mM CaCl $_2$ and infected with 4 x10⁵ pfu of the phage cDNA library for 15 minutes at 37°C. Infected cultures were then mixed with 10 ml

of NZC broth containing 1 % agarose at 55°C and 20 spread on plates containing NZC broth + 1.5 % bacto-agar in 150 mm petri dishes. Plates were incubated at 37°C overnight and then stored at 4°C. These plates were referred to as the master phage cDNA library. 25

Pre-cut 82 mm HAHY nitrocellulose filters (Millipore) were wetted in ${\rm H_2O}$, soaked briefly in 1 M NaCl and blotted dry on paper towels. Multiple plate lifts were made by placing wetted nitrocellulose

filters on top of each chilled master plate of the 30 phage cDNA library for 30 to 90 seconds. Filters were keyed to the plate by asymmetrical stabbing a 20 ga syringe needle containing india ink through the filter and into the agar plate. The filters were 35

then removed and phage DNA was fixed to the filters using the same procedure described above for lysis of bacterial colonies. The filters were then air-dried

	for 30-60 minutes and baked for 2 hours at 70°C	
	in vacuo. Pairs of filters were placed in	
5	heat-sealed bags with the plaque sides oriented	
	outwards and prehybridized with 6X SSC, 25 mM sodium	
	phosphate pH 6.8, 1 mM EDTA, 1 % SDS and 100 µg/ml	•
	sheared and denatured calf thymus DNA for 6-7 hours	,
	at 65°C with occasional mixing.	
10	Plasmid pIn 2-1 was nick-translated as	
	described above to a specific activity of 2.5 x 108	
	cpm/µg of DNA, and purified by spun-column	
	chromatrography using Sephadex® G-50. Prehybrid-	
	ization buffer was removed for	
15	ization buffer was removed from the bags containing	
	the replicas of the phage library and replaced with	
	20 ml of the same buffer containing 1.5 X 106 cpm of	
	denatured pIn 2-1 probe per ml of hybridization solution. Filters were hybridized at 6500	
20	with occasional mixing. Filters were removed from	
20	the bags, washed twice at room temperature for 15	
	minutes with 2X SSC, 0.5% SDS, and twice at 65°C for	
	30 minutes with 0.1% SSC, 0.1% SDS buffer. The	
	filters were briefly air dried, wrapped in	
25	polyethylene wrap and exposed to Kodak X-OMAT XAR-5	
25	film at -80°C overnight using a single Du Pont	
	Lightning Plus intensifying screen.	
	Plaques hybridizing with the pIn 2-1 probe were	
	picked from the master plates. Stocks of these	
20	hybridizing phage were made by removing agarose plugs	
30	from the plates containing appropriate planues	
	placing them in numbered 1.5 ml microfuge tubes	
	containing 1 ml of SM buffer with 1 drop of	
	chloroform and allowing the phage to diffuse out of	
	the plugs overnight at 4°C. Plaque purification was	
35	performed on each phage by serially diluting the	
	phage stocks, infecting 100 µl of an overnight	

culture E. coli Y1090 with 100 µl aliqouts of the dilutions and growing them on NZC plates as described above. Lifts of these plates were made and hybridized with a labelled $^{32}\mathrm{P}$ pIn 2-1 cDNA as previously described. Hybridizing plaques were repeatedly subjected to this procedure until all plaques on a given plate hybridized with the 2-1 cDNA 10 probe. Small scale phage DNA preparations of the $\lambda 2-1$ cDNA clones were made using the procedure in Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, N. Y., 15 (1982). The phage DNAs were digested to completion with EcoRI and analyzed by electrophoresis in a 1 % agarose gel. Results of this analysis showed one phage clone, designated 2-1.12, harbored a 900 bp insert. The insert contained a single internal Eco RI site that divided it into a 300 bp and a 600 bp 20 fragment when digested with Eco RI. This insert was of sufficient size to be a full length copy of the 2-1 mRNA. Restriction mapping of the pIn 2-1 and $\lambda 2-1.12$ cDNAs insert showed that $\lambda 2-1.12$ contained a 25 complete copy of the pIn 2-1 cDNA and that all missing 2-1 RNA sequences were probably present in λ2-1.12. The 600 bp Eco RI fragment from $\lambda 2-1.12$ was subcloned into the plasmid vector pUC18. To 30 accomplish this, pUC18 DNA was digested to completion with Eco RI. After digestion, a one-tenth volume of 1 M Tris-HC1 pH 8.4 was added directly to the tube. Calf intestinal alkaline phosphatase (CIAP) was then added using 0.5 units per microgram of DNA. The 35 dephosphorylation reaction was performed at 55°C for

30 minutes. CIAP was inactivated by sequential

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extractions of the DNA with equal volumes of phenol. phenol:chloroform (1:1 v/v) and chloroform. The DNA was then precipitated with ethanol in the presence of 0.25 M sodium acetate pH 6.0, collected by centrifugation and redissolved in TE, pH 8.0. $\lambda 2-1.12$ DNA was digested to completion with Eco RI, and equimolar aliquots of dephosphorylated, Eco RI digested pUC18 DNA and Eco RI digested $\lambda 2-1.12$ DNA were ligated together overnight at 16°C using ligation conditions described earlier. An aliquot of the ligation mixture was used to transform frozen competent E. coli HB101 cells (Bethesda Research Laboratories). Transformation of competent cells was accomplished by removing the cells from storage at -80°C and thawing them on ice. The ligation mixture was diluted 5 fold with H2O and an aliquot of this

- dilution was mixed with 100 µl of competent cells. The mixture was incubated on ice for 30 minutes and 20 then heat shocked for 45 seconds in a 42°C water bath without shaking. The cells were returned to ice for 2 minutes and diluted with 0.9 ml of S.O.C. medium (2% Bacto-tryptone, 0.5% yeast extract, 10 mM NaCl. 25
- 2.5 mM KCl, 10 mM MgCl2, 10 mM MgSO4 and 20 mM glucose). The cells were then shaken at 225 rpm at 37°C for 1 hour and aliquots of the transformation mixture were spread onto LB plates containing 50 µg/ml of ampicillin. Plates were then incubated at 30 37°C overnight. Small scale plasmid preparations
- were performed on individual amp-resistant colonies and aliquots of the DNA were digested with EcoRI until a colony was found that contained the 600 bp EcoRI fragment from pIn 2-1.12 ligated into pUC18. 35 This plasmid was called pIn2-1.12A.

DNA Sequence Analysis of 2-1 cDNA Clones

The nucleotide sequence of the 2-1 mRNA was determined by sequence analysis of pIn 2-1 and \(\lambda 2-1.12A\). The insert of pIn 2-1 was subcloned into the vector M13mp18 in order to perform dideoxy sequencing. For subcloning, an aliquot of pIn 2-1 was digested to completion with Pst I and the resulting 450 bp fragment was subcloned into the Pst I site of M13mp18 RF vector. An aliquot of the ligation mixture was used to transfect E. coli JM 101 and aliquots of the transfection reaction were plated on LB plates containing X-Gal and IPTG and grown

overnight at 37°C. Individual white plaques were analyzed until a phage was found that contained the CDNA insert in the Pat I site of M13. A DNA

sequencing template was prepared from this phage by 20 scooping a portion of a plaque out from the agar and using it to inoculate 3 ml of 2 X YT media in a 15 ml falcon tube containing 200 µl of expotentially growing JM 101 cells. The culture was incubated at 37°C with vigorous shaking for 5 hours. A 1 ml aliquot of the phage culture was removed and

centrifuged in a 1.5 ml microfuge tube for 5-10 minutes at 4°C. One ml of phage supernatant was carefully pipetted off and placed into a fresh tube containing 200 µl of 20% PEG 8000, 2.5 M NaCl. The tube was inverted several times, and then incubated

at room temperature for 20-30 minutes. The phage were collected by centrifugation for 10 minutes in a microfuge at room temperature. The supernatant was carefully removed and the tube was recentrifuged to

35 remove any remaining superantant from the tube walls. The phage were resuspended in 100 μl of 10 mM

Tris-HCl, pH 7.6 and extracted with 50 µl of phenol:chloroform (1:1 v/v) by vortexing the tube. The tube was centrifuged for 5 minutes at room temperature and the upper aqueous phase was transferred to a next tube.

temperature and the upper aqueous phase was transferred to a new tube. Phage DNA was precipitated with 25 µl of 2 M sodium acetate, pH 7.0 and 320 µl of ethanol at -70°C for 10 minutes or

10 overnight freezing at -20°C. The DNA, suitable for use as a sequencing template, was collected by centrifugation in a microfuge at 4°C for 10-20 minutes and dissolved in TE pH 8.0.

This template DNA was sequenced using the dideoxy method of Sanger [Sanger, F. et al., Proc. Natl. Acad. Sci USA, 74:5463, 1977], using a dideoxy sequencing kit (Pharmacia Inc., 800 Centennial Avenua, Piscataway, NJ 08854, CAT # 27-1555-01)

following the manufacturer's recommended procedures.

A portion of the 2-1 DNA insert in the M13
clone was deleted by cutting the RF DNA with Eco RI
and religating the DNA back together. This removed
approximately 170 bp from the cDNA insert adjascent
to the sequencing primer in the vector. This

Subclone was sequenced.

25 subclone was sequenced as above using the universal primer to complete the sequencing of the pIn 2-1 CDNA clone.

The cDNA clone, 2-1.12A, was sequenced to complete the sequence of the 2-1 mRNA. The pIn 30 2-1.12A sequence was determined by the method of Maxem and Gilbert (Maxem, A.M. and Gilbert, W., Methods in Enzymology, 65:499-512, 1980) with modifications described by Barker et al. (Barker et

al., Plant Molecular Biology, 2:335-350, 1983). DNA Sequence analysis confirmed the identity of pln 2-1 and pln 2-1.12 since 200 bp region common to both clones shared an identical nucleotide sequence.

Isolation of 2-1 Genomic Clone 21.14

Plant material used for DNA isolation was

obtained from greenhouse grown plants of the inbred
corn line Missouri 17 (Mo17). Leaf material from
vegetative plants was harvested, deribbed, and frozen
in liquid nitrogen. High molecular weight DNA was
isolated from 30 g of leaf material as follows:

15 frozen leaf material was placed in a coffee grinder
along with a small amount of dry ice and ground to a
fine powder. After the dry ice had sublimed, the
frozen powder was transferred to a beaker and

- suspended in 100 ml of ccld buffer A (100 mM Tris-HCl 20 pH 9.0, 100 mM NaCl, 10 mM MgCl₂, 0.5 M sucrose, 0.1% β-mercaptoethanol, 0.4% diethylthiocarbamic acid). Nuclei were pelleted from the slurry by centrifugation at 10,000 rpm for two minutes in a Sorvall GSA rotor. The supernatant was discarded and 25 the nellet was resuspense.
- 25 the pellet was resuspended in 3 ml of buffer A. The nuclei were lysed by resuspending them in 20 ml of lysis buffer (100 mM Tris-HC1 pH 8.3, 100 mM NaCl, 50 mM Na_ZEDTA, 1.5 % SDE, 15% phenol) and incubating the mixture at 55°C for 10 minutes with constant
- ostirring. Ten ml of 5 M potasium acetate was then added and the mixture was placed on ice for 10 minutes to precipitate SDS, SDS-protein complexes and SDS-cell wall complexes. The precipitate was collected by centrifugation at 5000 rpm for 10 min.
- 35 in a Sorvall table-top centrifuge. The supernatant was transferred to a new tube, and the solution

	nes extracted with an equal volume of	
	chloroform:isoamyl alcohol (24:1 v/v) after page:	
5	or 3 mi or 10 M ammonium acetate . DNA was about	,
	precipitated by addition of an equal volume of	
	isopropanol, collected by centrifugation and	_
	resuspended in 30 ml H ₂ O. Solid cesium chloride was	•
	added using 0.9 g for each ml of solution and	
10	ethidium bromide was added to 300 μg/ml. DNA was	
	Centrifuged at 45,000 rpm for 16 hours in a Beckman	
	VTi50 rotor. Banded DNA was recovered from the	
	gradient by side puncturing the centrifuge tubes	
	with a 16 gauge needle and removing the band. The	
15	DNA was diluted to 30 ml with 1 g/ml CsCl (prepared	
	by adding 100 g CsCl to 100 ml TE pH 8.0) and banded	
•	once again following the same procedure. Ethidium	
	bromide was removed from the DNA by repeated	
	extractions with sodium chloride-saturated,	
20	water-saturated isopropanol. The DNA was then	
	precipitated with isopropanol. Mol7 genomic DNA was	
	collected by centrifugation and resuspended in TE pH	
	8.0.	
	An Mol7 genomic library was constructed as	
25	rollows: 100 micrograms of Mol7 DNA ware district	
	with 24 units of restriction enzyme Sau 3A in Cutsall	
	(100 mm potassium chloride, 20 mm Tris-HCl pu 7 r	
	may p-mercaptoethanol, 7 mM magnesium chloride)	
	Titth of the reaction was removed after 2 4 6 5	
30	and 10 minutes of digestion and the reaction was	
	scopped by adding EDTA to 50 mM. The five him-	
	points were pooled, extracted with an equal makes	
	phenol: entorororm; isoamyl alcohol (25:24.)	•
	and DNA in the pool was ethanol precipitated	
35	collected by centrifugation. The DNA was discalled	•
	In V.1 mi H2O and loaded on a 10-40% glygorol	
	gradient (10-40% glycerol in 1 M Nacl 20 - m m :	
	pH 8.0, 1 mM EDTA). Centrifugation was performed at	

40,000 rpm for 16 hours in a Beckman SW 41 rotor. Fractions (0.4 ml) were collected from the bottom of 5 the polyallomer tube through a wide bore needle and aliquots of the fractions were analyzed by electrophoresis in a 0.9% agarose gel. Fractions containing 12-20 kbp DNA fragments were pooled, extracted with an equal volume of phenol/chloroform 10 (1:1 v/v), precipitated with ethanol and resuspended in TE pH 8.0. Four-tenths of a microgram of this size-fractionated DNA was ligated overnight to 1 microgram of Eco RI-Bam HI digested lambda EMBL 3 DNA (Stratagene) using 5 weiss units of DNA ligase (New England Biolabs) in ligase buffer (50 mM Tris-HCl pH 15 8.0. 10 mM dithiothreitol, 10 mM magnesium chloride, 1 mM ATP) at 15°C for 24 hours. Ligated DNA was packaged using Gigapack Gold packaging extracts

(Stratagene) following the manufacturer's recomended
20 protocol.

A library of 500,000 phage was plated on 150 mm diameter LAM plates (10 g Bacto-Tryptone, 5 g yeast extract, 10 g NaCl, 2.5 g MgSO4.7H20, 10 g agarose per liter, 80 ml per plate) at a density of 25 about 25,000 plaques per plate. To do this, phage (in a volume of less than 200 μ 1) were added to 200 μl of 10 $\dot{m}M$ $CaCl_{2}$, 10 mM $MgCl_{2}$ and 200 μl of an overnight E. coli LE 392 culture grown in 2XYT (16 g Bacto-tryptone, 10 g yeast extract, 5 g NaCl 0.2% maltose, water to 1 liter) and phage were allowed to 30 adsorb to host cells at 37°C for 10-15 minutes. This culture was then added to 8 ml molten 0.8 % top agarose (10 g Bacto tryptone, 2.5 g NaCl, 0.2 g MgCl₂, 8 g agarose, water to 1 liter) at 50°C and poured onto LAM plates. After the top agarose 35 hardened, plates were incubated at 37°C overnight.

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	Phage lifts were performed the next morning by
	laying dry mitrocellulose filters (Millipore) on the
5	surface of the plates for 5 minutes. Filters were
	then transferred to a piece of Whatman 3MM paper that
	was saturated with 0.5 M NaOH, 1.5 M NaCl. After 5
	minutes the filters were transferred to a sheet of
	3MM paper saturated with 0.5 M Tris-HCl pH 7.5, 1.5 P
10	NaCl. After 5 minutes the filters were transferred
	to a piece of 3MM paper saturated with 2X SSC for
	5-10 minutes. The filters were then baked at 80°C
	for two hours in vacuo.
	Filters were prehybridized at 42°C for 4 hours
15	in a 150 mm glass crystallizing dish using 150 ml of
	prehybridization buffer (50% deionized formamide, 5%
	SSC, 100 µg/ml denatured salmon sperm DNA, 0.05% SDS,
	0.05 M sodium phosphate pH, 0.1% Ficoll, 0.1%
	polyvinylpyrolidine, 0.1% BSA). One µg of plasmid
20	DID 2-1 was night become total to so a second

10 pln 2-1 was nick translated in 50 µl of 50 mM Tris-HCl pH 7.2, 10 mM MgSO₄, 0.1 mM DTT, 50 mg/ml BSA, 10 uci 32P GATP (Amersham), 2 µg/ml DNsse (Sigma DN-EP), 20 uM dATP, dTTP, dGTP, 5 units DNA polymersse I (BMB) at 15°C for 1 hour. The reaction

25 was stopped by adding 1 µl of 0.5M EDTA, and DNA was then precipitated by adding 50 µl of water, 30 µl of 10 M ammonium acetate, 10 µg yeast tRNA carrier and 350 µl of ethanol. The DNA was collected by centrifugation, dissolved in 0.5 ml H₂O, and

denatured by heating for 5 minutes in a boiling water bath followed by quick cooling on ice. Prehybridization solution was discarded and the filters were probed overnight at 42°C with

nick-translated pIn 2-1 with hybridization buffer 35 (50% deioniz d formamide, 5% SSC, 100 µg/ml denatured salmon sperm DNA, 0.05% SDS, 0.02 M sodium phosphate 0.2% Ficoll, 0.02% Bowine serum albumin, 10% dextran sulfate) using 5 x 10⁵ cpm per ml of buffer. The next morning, filters were washed twice for 20 minuses.

5 buffer. The next morning, filters were washed twice for 20 minutes in 1X SSC, 0.5% SDS at room temperature and three times for 20 minutes in 0.1X SSC, 0.5% SDS at 65°C in a shaking water bath. The filters were blotted dry between two sheets of 3MM Paper, wrapped in polyethylene food wrap, and exposed on Kodak XAR-5 film overnight at -80°C using a single Du Pont Lighting Plus intensifying screen. Films were developed using a Kodak X-OMAT developer.

Positive plaques were picked by taking plugs
from the agar plates with the thick end of a pasteur
pipette and placing them in 0.5 ml of SM. Dilutions
of the phage in each plug were used to infect E. coli
LE 392 as before and plated on 80 mm diameter LAM
plates using 3 ml top agarose, 100 µl 10 mM CaCl₂, 10

20 mM MgCl₂, and 100 µl of an overnight LE392 culture. Purification was performed on each phage plaque as described earlier. The phage lift-pick-plating cycle was carried out until pure plaques were obtained. Fifteen pure phage isolates, designated 21.1 to 21.15 were grown in liquid culture for isolation of DNA.

25 were grown in liquid culture for isolation of DNA. Single pure plaques were removed from plates and eluted into 0.5 ml of SM. Fifty µl of these phage stocks were incubated with 50 µl of a two times concentrated overnight LE392 culture in 10 mM MgCl₂

o at 37°C for 15 minutes. The infected bacteria were then added to 20 ml of pre-warmed LB (10 g Bacto-Tryptone, 5 g Bacto yeast extract, 5 g NaCl, 1 g glucose, water to 1 liter) and shaken at 37°C, 180-200rpm. The cultures generally lysed after 4-7

35 hours. Chloroform was added to a concentration of 1%, and the lysates were shaken for an additional 10 minutes. Cellular debris was removed by

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centrifugation at 10,000 rpm for 10 minutes in a Sorvall SS 34 rotor. The supernatants were transferred to new tubes and DNAse I and RNAse A were added to 20 µg/ml and 10 µg/ml respectively. After a 15-30 minute incubation at 37°C, phage were precipitated by addition of a one fifth volume of 20% PEG 8000, 2.5M NaCl to the lysate. After 15 minutes at room temperature, the phage were collected by centrifugation at 15,000 rpm for 15 minutes at 4°C in a Sorvall SS-34 rotor and resuspended in 0.5 ml of SM. Fifty µl of 0.5 M EDTA, 70 µl of 10 % SDS and 300 ul of phenol were added to phage suspensions to lyse them. The lysates were extracted with phenol:chloroform (1:1 v/v), and DNA in the aqueous phases was precipitated by adding one-tenth volume 3.0 M sodium acetate and two-thirds volume

isopropanol. DNA was collected by centrifugation and the pellet was washed with 70% ethanol, dried and resuspended in 50 μH_2O .

Identification and Characterization of Genomic Clone 21.14

The fifteen genomic clones were first characterized by restriction mapping in an attempt to find regions in the clones corresponding to the 2-1 cDNA. Two μg of DNA was digested with several different restriction enzymes in ten μl of 1X cutsall (or 1.5 x cutsall for Sal I) and analyzed by electrophoresis using 1% agarose gels. Restriction maps generated for each of the clones failed to identify candidate genes for further analysis. Therefore, these genomic clones were mapped using a

35 probe made by randomly primed cDNA synthesis using

RNA from N-(aminocarbonyl)-2-chlorobenzenesulfonamidetreated corn roots as a template to identify regions in the various genomic clones that corresponded to the coding region of the genes. Phage DNA was digested with a variety of different restriction enzymes and the digestion products were separated by electrophoresis using 1% agarose gels. The DNA was 10 transferred to Gene Screen Plus membranes (New England Nuclear) and hybridized with a randomly primed cDNA probe that was made as follows: 1 ug poly(A) + RNA from N-(aminocarbony1)-2-chlorobenzenesulfonamide-treated corn roots was added to 30 ul of water and placed in a boiling water bath for 5 15 minutes. After cooling on ice, 10 µl of 10X first strand buffer (0.5 M Tris-HCl pH 8.5, 0.4 M KCl, 0.1 M MgCl₂, 0.4 mM DTT,) 2.5 μl 2 mM dATP, 2.5 μl 2 mM dCTP, 5 µl 20 mM dGTP, 5 µl 20 mM dTTP, 1 µl RNAsin 20 (Promega Biotech, Inc.), 20 µl random hexamer primers (16 µg/ul, Pharmacia, cat #272266-01 or P-L Bichemicals, cat #PLB9223), 10 μl α32P dATP (100u Ci), 10 μ l α 32P dCTP (100 μ Ci), and 20 units of reverse transcriptase were added. The reaction mixture was then incubated at 37°C for one hour. 25 reaction was stopped by adding 10 μ 1 0.5 M EDTA. was hydrolyzed by adding 50 μ l 0.15 M NaOH and heating the mixture for one hour at 65°C. Base was then neutralized by adding 25 μl of 2 M Tris-HCl pH 30 8.0 and 50 μ l 1M HCl. The DNA was precipitated with ammonium acetate and ethanol in the presence of carrier tRNA as described earlier. The randomly primed cDNA probe was then dissolved in 0.5 ml of H₂O. Hybridization and prehybridizations were carried out as described above for genomic library 35

screening. Data from this restriction mapping

	analysis defined areas of each genomic clone that
	were homologous to pIn 2-1, but failed to identify
5	any genomic clones corresponding to the In2-1 cDNA.
	Therefore, restriction fragments of the genomic
	clones that hybridized to the random cDNA probe were
	subcloned into either pUC19 or the vector Bluescript
	pBS(+), (Stratagene) for DNA sequence analysis.
10	Subcloning of genomic DNAs were performed by
	digesting 10 µg of phage DNA and a suitable vector
	(either pUC19 or pBS+) with the appropriate
	restriction enzymes. The DNAs were extracted with
	phenol/chloroform (1:1 v/v), precipitated with
15	ethanol and resuspended in 10 µl of TE. Phage DNA
	was ligated to vector DNA in a final volume of 10
	μl. After an overnight incubation at 15°C, the
	ligation products were used to transform competent
	JM83 cells. Colonies harboring the desired plasmids
20	were identified by performing small scale plasmid
	preparations and digesting aliquots of the resulting
	plasmids with diagnostic restriction enzymes.
	The strategy used to sequence subcloned genomic
	fragments was to create a nested set of deletions for
25	each subclone using Bal 31 nuclease (New England
	Biolabs). Plasmid DNA (20 µg) was linearized using
	an appropriate restriction enzyme and then extracted
	once with phenol:chloroform (1:1 v/v) and
	precipitated with ethanol. DNA was collected by
30	centrifugation, washed once with 70% ethanol, dried,
	and resuspended in 100 μl of H_2O . Nuclease digestion
	was carried out in a total volume of 250 µl using 20
	units Bal 31 under the assay conditions described by
	the manufacturer. Aliquots of 10 µl were removed at
35	various times ranging up to 8 minutes and pooled into
	5 groups. The reactions were stopped by adding the

	Phage lifts were next performed as described
	above. Lifts were probed with 32P-labelled gel
5	purified-insert from the plasmid on which deletions
	were performed to detect plaques with Bal 31
	deletions. Plaques hybridizing to the probe were
	picked and grown as follows: a positive plaque was
	stabbed with a sterile toothpick which was then put
10	into 2 ml 2XYT containing 10 µl of a JM101 overnight
	culture. The culture was grown for 5 hours at 37°C
	and small scale plasmid preparations were performed.
	One ml of overnight culture was poured into a
	microfuge tube and centrifuged for 20 seconds. The
15	supernatant was poured off into a new tube and saved
	for later preparation of single-stranded DNA. The
	pelleted cells were resuspended in 0.35 ml of BPB (8%
	Sucrose, 0.5% Triton X-100, 50 mM EDTA pH 8.0, 10 mM
	Tris-HCl pH 8.0). Twenty five µl of a freshly
20	prepared lysozyme solution (10 mg/ml in BPB) was
	added and the tube was placed in a boiling water
	bath for 40 seconds, followed immediately by
	centrifugation for 10 minutes at room temperature in
	a microcentrifuge. Chromosomal DNA as well as other
25	debris formed a gelatinous pellet, and was removed
	with a sterile toothpick. Plasmid DNA was
	precipitated by addition of 30 µl 3M sodium acetate
	and 250 µl isopropanol. DNA was recovered by
	centrifugation, washed with 70% ethanol, dried and
30	resuspended in 75 µ1 H20. Six µ1 aliquots were
	digested (in Cutsall) with appropriate enzymes that
	would excise the inserts. After analysis of these
	digestions by electrophoresis on 1% agarose gels, the
	subclones were ordered in decreasing order f size
35	(increasing amount of Bal 31 deletion) and clones
	were chosen so that a series of progressive 100 bp
	deletions of the starting clone subjected to poly

deletion was obtained.

Single stranded DNA for dideoxy chain-

- termination sequencing was isolated from the 1 ml of 5 the supernatant saved at the start of the small scale plasmid preparation procedure. The supernatant was mixed with 150 µl 20% PEG 8000, 2.5 M NaCl and phage were collected by centrifugation for 5 minutes in a microcentrifuge after 15 minutes at room
- 10 temperature. All traces of supernatant were removed by aspiration, and the pellet was resuspended in 100 µl 0.3 M sodium acetate, 1 mM EDTA. Phage were lysed by extraction with an equal volume of phenol:chloroform (1:1 v/v) and DNA was precipitated with
- 15 ethanol. DNA was collected by centrifugation, washed with 70% ethanol, dried briefly and resuspended in 25 µl of H₂O.

Sequencing was performed using the M13 universal -10 17mer primer (New England Biolabs,

- 20 Inc.) The annealing reaction was performed at 60-65°C for 1 hour using 3.5 µl template DNA, 2.5 µl annealing buffer (100 mM Tris-HCl pH 8.5, 50 mM MgCl₂), 1 µl universal sequencing primer (1 ng/ul) and 4 µl water. The annealed DNA was then placed on ice. The components of the sequencing reaction
- were: 1) Termination mixes containing dideoxy A.C.G, or T plus deoxy A.C.G, and T in appropriate ratios; 2) polymerase cocktail which contained 0.9 µl 0.1 M Tris-HCl pH 8.3, 1 µl (10u Ci) 355 dATP, 1 µl 0.1M 30 dithiothreitol, 6.1 µl water, 0.25 µl Klenow (5
- units/ul). Two µl of each component were mixed in a well of a microtiter dish 4 wells (A,C,G,T) for each annealing and incubated at 37°C for 20 minutes. At this time, 2 µl of chase solution (a
- 35 solution containing 0.5 mM of all four dNTPs) was added to each well. After an additional 25 minutes of incubation, terminating dye (0.08 % bromphenol blue, 0.08 % xylene cyanol, 20 mm EDTA in deionized

formamide) was added to the wells. The reactions were heated uncovered at 90°C in an oven for 10 5 minutes, placed on ice, and subjected to electrophoresis in a 6 % polyacrylamide gel in 1X TRE (0.089 M Tris-borate, 0.089 M boric acid, 0.002 M EDTA) containing 8 M urea at 1500 volts for approximately 2 hours. Urea was removed from the gel by soaking it in 10% methanol, 10% acetic acid for 15 10 minutes. The gel was then transferred to a sheet of Whatman 3MM paper and dried on a gel dryer with vacuum. The gel was autoradiographed with Kodak X-AR film overnight at room temperature with no 15 intensifying screen. DNA sequences were read from the gel, entered into a computer and analyzed using the Cold Spring Harbor programs. The sequence of the promoter from the 21.14 gene extending 5' from the Nco I site that initiates protein synthesis is shown 20 in Figure 2. Sequence analysis also revealed that a 1.9 kbp Eco RI/Sal I subclone of genomic clone 21.14 contained sequences for the 3' half of 2-1 mRNA. This subclone was designated pJE482-62. Similarly, a 4.8 kbp Eco RI/Sal I subclone from 21.14 was shown to 25 contain the coding sequences for the 5' half of the 2-1 mRNA. This clone was designated pJE 484-1. Complete sequence analysis revealed that genomic clone 21.14 contained a perfect copy of the 2-1 coding sequence distributed among 9 exons and eight 30 introns. Therefore 21.14 was designated as a gene

Cloning and Mutagenesis of the Regulatory Regions of the 21.14

encoding the 2-1 mRNA

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After identifying a genomic clone whose sequence agreed perfectly with that of the 2-1 cDNA clones, a search was begun for the regulatory

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regions of the gene. The first codon iniating

- 5 the 21.14 gene by its position and homology to the consensus sequence A..ATGG, as well as by comparison of the 21.14 genomic sequence to the open reading frame in the 2-1 cDNA sequence.
- 10 Construction of plasmids p484-1(Nco I) and p484-62 (Bg1 II)

Site directed mutatgenesis was performed on the regulatory regions of the 21.14 genomic clone so that the expression of a foreign coding sequence could easily be placed under the control of chemicals known to affect the expression of the 2-1 gene. An

oligonucleotide of the sequence
5'-GAGCTGCGGTACCGGC-3' was designed to introduce an

Nco I restriction site in pJE484-1 at the ATG codon corresponding to the start of the 2-1 protein coding region of the message. Another oligonucleotide, 5'-TGAGATCTGACAAA-3', was designed to introduce a Bg1 II restriction site in pJE482-62 at the 3' end of the gene, 9 base pairs past the termination codon of the 2-1 protein. Both oligonucleotides were synthesized using an Applied Biosystems DNA synthesizer.

The plasmid pJE 484-1 was transformed into the dut ung E. coli. strain BW313 [disclosed in Proc. Natl. Aacd. Sci., USA, Vol. 79, pp 488-492 [1982]]. Cultures were grown for the production of

single-stranded DNA as described earlier in this Example. Colonies were picked with sterile toothpicks and used to inoculate two 5 ml tubes of 2XYT containing 100 µg/ml ampicillin and 5 ul of a

35 M13K07 stock (a helper phage for packaging of single-stranded DNA; titer 10¹¹ pfu/m1). The

	cultures were shaken at 37°C, and after two hours of
	growth, kanamycin was added to 50 μg/ml. The
5	incubation was then continued at 37°C overnight. The
	tubes were pooled and bacteria were removed by
	centrifugation at 8,000 rpm for 10 minutes in a
	Sorvall HB4 rotor at 4°C. Six ml of the supernatant
	were removed to a new tube and 1.5 ml 20% PEG, 2.5 M
10	NaCl added to it and mixed well. After 15 minutes at
	room temperature, phage particles were pelleted by
	centrifugation at 8,000 rpm for 10 minutes in a
	Sorvall HB4 rotor. The pellet was resuspended in 0.4
	ml 2XYT and transferred to a new microfuge tube.
15	Phage particles were precipitated by adding O.1 ml
	20% PEG, 2.5 M NaCl. After 5 minutes, phage were
	collected by centrifugation and all traces of
	supernatant were removed by aspiration. Phage
	particles were resuspended in 0.5 ml 0.3 M sodium
20	acetate, 1 mM EDTA, and extracted with .
	phenol:chloroform (1:1 v/v). Phage DNA was then
	precipitated with ethanol, collected by
	centrifugation, washed once with 70% ethanol and
	resuspended in 50 µl of H2O. The concentration of
25	DNA was determined by measuring the absorbance of a 1
	to 50 dilution of this solution. 0.5 pmole of this
	single-stranded DNA was annealed to 25 pmole of the
	oligonucleotide 5'-GAGCTGCGGTACCGGC-3' in 20 µl Fritz
	standard annealing buffer (8X annealing buffer is 1.5
30	M KCl, 100 mM Tris-HCl pH 7.5) for 30 minutes at
	55°C, 15 minutes at 37°C, and 15 minutes at room
	temperature. After annealing, 2.3 µl 10% fill-in
	buffer (0.625 M KC1, 0.275 M Tris-HC1 pH 7.5, 0.15 M
	MgCl2, 20 mM DTT, 2 mM ATP, 1 mM of each dNTPs), 1 ul

35 Klenow (5 U/µl) and 1 µl of 0.6 U/µl DNA ligase were

	added. The tube was incubated overnight at room
	temperature. The next day, competent E. coli strain
5	JM83 was transformed with the products of this
	ligation reaction (as described earlier) and plated
	on LB plates containing 100 µg/ml ampicillin. Small
	scale plasmid preparations were performed on the
	resulting colonies and the DNA was digested with Nco
10	I until a transformant was found that contained a
	plasmid that was linearized by Nco I, indicating that
	the desired mutation had taken place. This new
	plasmid was designated pJE 484-1(NcoI) (Figure 3). In
	the same manner, the plasmid pJE 484-62 was
15	mutagenized with the oligonucleotide
	5'-TGAGATCTGACAAA-3' to create a new Bgl II site
	downstream of the translation stop site of the 2-1
	protein. This new plasmid was designated pJE
	484-62(Pg) TT) (Pieure 2)

Identification of the Transcription Start Site of 21.14 Gene

Primer extension analysis was performed to determine the transcription start site of the 21.14 25 gene using a method based on the procedure of McKnight [McKnight, S. L., Cell 31 355-366 1982]. A synthetic oligonucleotide, designated HH17, which is the reverse complement to bases 572 to 593 of the coding strand of the 21.14 gene (Figure 1) was synthesized using an Applied Biosystems Model 380A 30 DNA synthesizer. The HH17 oligonucleotide, 5'-CATGTCGTCGAGATGGGACTGG-3', was end-labeled with 32P-gamma ATP (specific activity 3000 Ci/mole, NEN Research Products) as follows: 5 µl (8.34 pmoles) of 35 32P-gamma ATP was dried in a microfuge tube in vacuo. The pellet was dissolved in 2 µl of HH17

primer (5 pmole) and 2 µl of 2.5% kinase buffer (1% buffer is 50 mM Tris-HCl pH 9.5, 10 mM MgCl2, 5 mM 5 dithiothreitol, 1 mM spermidine, 0.1 mM EDTA). One ul of T4 polynucleotide kinase (5.3 U/ul, Pharmacia) was added, and the labeling was allowed to proceed at 37°C for 15 minutes. The reaction was stopped by adding 75 ul TE (10 mM Tris-HCl pH 8, 1 mM EDTA), 54 μl 5 M ammonium acetate, 20 μg yeast tRNA carrier and 10 350 µl ice-cold ethanol. The oligonucleotide was precipitated on dry ice for 30 minutes and recovered by centrifugation at 4°C. The pellet was dissolved in 90 µl of TE pH 8.0 and re-precipitated on dry-ice 15 for 30 minutes after adding 10 µl 3 M sodium acetate. pH 6 and 250 ul ice-cold ethanol. The oligonucleotide pellet was collected as before. rinsed with 95% cold ethanol and dried in vacuo. The pellet was dissolved in 50 µl of 10 mM Tris-HC1 pH 8 20 at a final concentration of 0.1 pmole/ul and stored a+ 4°C Eight µg of total RNA isolated from the roots

of Mol7 corn plants treated hydroponically for 6 hours with 200 mg/l N-(aminocarbonyl)-2-chlorobenzenesulfonamide was mixed with 2 μ l (0.2 pmole) 25 32p-labeled HH17 primer, 2 µl of 5X annealing buffer (1.25 M KC1, 10 mM Tris pH 7.9) and 1 μ l of 30 mM vanadyl ribonucleoside complex (Bethesda Research Labs) at 0°C. Annealing was performed by heating the 30 mixture at 65°C for 3 minutes and cooling to 35°C over a 2 hour period. Primer extension was performed by adding 23µl of PE mix (10 mM MgCl2, 5 mM dithiothreitol, 20 mM Tris HCl pH 8.3, 0.33 mM of each dATP, dCTP, dGTP, dTTP, 100 µg/ml actinomycin-D), 0.5 µl of AMV rev rse transcriptase 35

(10 U/ul. Molecular Genetic Resources) to the tube

followed by incubation at 37°C for 45 minutes.

Primer extension products were precipitated on dry

- 5 ice for 20 minutes after adding 300 µl of ice-cold ethanol. The precipitate was collected by cenrifugation at 4°C, rinsed with 70% ice-cold ethanol and dried in vacuo.
- The HH17 oligonucleotide was used as a primer for sequencing of plasmid pJE516 (described in Exemple 6). Four µg of pJE516 was denatured in 200 mM NaOH, 0.2 mM EDTA at room temperature for 5 minutes and base was neutrelized with 2 M ammonium
- acetate pH 5.4 at 0°C. The denatured DNA was
 precipitated on dry ice for 10 minutes after adding 2
 volumes of ice-cold ethanol. The DNA was collected
 by centrifugation at 4°C for 15 minutes, rinsed with
 70% ethanol, dried in vacuo and dissolved in 10 µl of
 water. Seven µl of denatured pUES16 was sequenced
- 20 with HH17 as the primer using a Sequenase® Rit (United States Biochemical Corporation) using the procedures recommended by the manufacturer.

The primer extension products from above were dissolved in 3 µl of 0.1 M NaOH, 1 mM EDTA for 30 minutes at room temperature. Six µl of gel loading buffer was then added and the solution was heated at 90°C for 5 minutes. Primer extension products and primed pJE516 sequencing reactions were separated by electrophoresis on a 12% polyacrylamide gel in 1X TBE 30 Containing 7 M urea. The gel was then dried and

autoradiographed. Analysis of the primer extension products showed the presence of one major band whose length corresponded to a transcription start site at base 532 of the 21.14 gene promoter fragment in

35 Figure 2 and two minor products corresponding to bases 533 and 536. The positions of these bases in

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3.0

Figure 2 is indicated by arrows. Nucleotide 532 of

2-1 corn gene promoter designated 21.14 was therefore
assigned as the major transcription start site.

EXAMPLE 2

Identification and Isolation of the Promoter and 3'

10 Downstream Regions of the 2-2#4 Corn 2-2 Gene

Isolation and Characterization of 2-2 cDNA clones
Details of the techniques used to perform the procedures used in Example 2 are presented in

Example 1. The cDNA library made using poly(A)* RNA from the roots of N-(aminocarbonyl)-2-chlorobenzenesulfonamide-treated Missouri 17 corn plants (described in Example 1) was analyzed for additional

cDNA clones representing mRNAs induced by substituted

benzenesulfonamides. The library was subjected to
differential screening as before and a new colony

displaying stronger hybridization with the cDNA probe made using RNA from roots treated with N-(aminocarbony1)-2-chlorobenzenesulfonamide was identified. This colony was designated In 2-2

A small scale plasmid preparation was performed on the plasmid contained in colony In 2-2. This plasmid was designated pIn 2-2. An aliquot of pIn 2-2 was nick translated as described earlier. A slot

blot containing total RNA from N-(aminocarbony1)-2-

chlorobenzenesulfonamide-treated and untreated roots was prepared and probed with nick-translated pIn 2-2 as described for pIn 2-1 in Example 1. This analysis confirmed that pIn 2-2 contained a cDNA insert that

35 hybridizes strongly to RNA from roots of plants

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treated with N-(aminocarbony1)-2-chlorobenzenesulfonamide, but not to RNA from untreated roots.

The pIn 2-2 small scale plasmid preparation was digested to completion with Pst I and analyzed by agarose gel electrophoresis. The cDNA insert of the plasmid was excised by Pst I as a single 1200 bp fragment. Nick-translated pIn 2-2 was used to probe a Northern blot of RNA from both untreated and N-(aminocarbonyl)-2-chlorobenzenesulfonamide-treated roots. This probe hybridized to a single 1.35 knt mRNA that was present only in RNA from the roots of N-(aminocarbonv1)-2-chlorobenzenesulfonamide-treated 15

plants. This indicated that the insert in pln 2-2 was not a full lenght copy of the 2-2 mRNA.

A new plasmid cDNA library was made to isolate full length 2-2 cDNA clones. An aliqout of the ds cDNA used to make the Agtll library described in

20 Example 1 was ligated overnight into the vector pUC18 that had been cut to completion with Eco RI and dephosphorylated. Aliquots of the ligation reaction were used to transform competent E. coli DH5 (Bethesda Research Laboratories) using the protocol 25 suggested by the manufacturer. A set of master

filters of this library was made by arraying individual ampicillin resistant colonies onto nitrocellulose as described for the plasmid cDNA library in Example 1. Another set of master filters were prepared by transferring colonies directly to 30 nitrocelluluse by laying a dry filter onto a plate

that contained 150-250 transformed colonies per plate. The filter was then removed and placed colony side up on a fresh LB/amp plate. Three replica 35

nitrocellulose filter copies of the library were pr pared and the DNA in each colony was fixed to the WO 90/11361 PCT/US90/01210

	84	
	filters as described previously. One set of replica	
_	filters was prehybridized and then hybridized with a	
5	mixed probe consisting of nick-translated plasmid	
	DNAs from a number of sources including the plasmid	•
	pIn 2-2. Plasmid nick-translation, and filter	
	prehybridization and hybridization were performed as	
	described for the identification of specific cDNA	
10	clones from the Agtll cDNA library in Example 1. A	
	total of 1500 colonies were screened, and twelve of	
	these colonies hybridized to the mixed cDNA probe.	
	These putative positive clones were	
	characterized by performing small scale plasmid DNA	
15	preparations from each colony. Plasmids were	
	digested to completion with Eco RI and the digestion	
	products were separated by agarose gel	
	electrophoresis. The DNA fragments in the gel were	
20	blotted to a Zeta Probe® membrane, and the blot was	
20	then hybridized with nick-translated pIn 2-2 to	
	identify the 2-2 clones in the mixed population, as	
	well as to obtain a size estimate for the insert	
	sizes of any new 2-2 clones that were found. Five	
	colonies hybridized to 2-2 probe, with one appearing	
25	to contain a full length 1.35 kbp insert. This clone	

Isolation of genomic clone 2-2 #4

was designated pIn 2-2-3.

The library of Mo17 genomic DNA used to obtain
genomic clones corresponding to the 2-1 cDNA was
screened for 2-2 genomic clones as described in
Example 1. Three 2-2 genomic clones were identified
and plaque purified from this library. The three
clones were mapped using a probe made by rendomly
primed cDNA synthesis using RNA from N-(amino
carbonyl)-2-chlorobenzenesulfonamide-treated corn

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roots as described in Example 1. The result of this analysis indicated that the clone designated 2-2 #4 contained a region of homology to the randomly primed CDNA probe in the center of its insert, and was therefore chosen for further analysis.

DNA Sequence Analysis of In2-2-3 and 2-2 #4 clones

A plate stock of phage 2-2 #4 was prepared by diluting 100 μ l of an overnight culture of LE392 with an equal volume of 10 mM MgCl2. 10 mM CaCl2. The diluted culture was incubated at 37°C for 20 minutes with 40 μ l of plaque purified 2-2 #4 phage.

- The culture was mixed with 3 ml of molten 55°C top agarose (0.7 % agarose in MZC broth), spread over the surface of a 100 mm NZC agar plate and grown at 37°C for 8 hours. The surface of the plate was covered
- with 6 ml of SM and it was placed at 4°C overnight on 20 an orbital shaker at 50 rpm. The SM was removed from the plate, mixed with 50 µl of CHCl₃, and stored at 4°C. Serial dilutions of this stock were titered on E. coli LE 392 to determine phage liter.
- A large scale preparation of genomic clone z-2

 #4 DNA was performed by diluting 3 ml of an overnight
 culture of E. coli LE392 grown in NZC medium with 3
 ml of 10 mM MgCl₂, 10 mM CaCl₂ and inoculating the
 bacteria with 2 X 10⁶ plague forming units (pfu) of
 2-2 #4. This culture was incubated at 37°C for 15-20
 minutes and then used to inoculate 500 ml of NZC at
 37°C. The culture was grown at 37°C with vigorous
 sgitation until lysis occurred (approximately seven
 hours). The lysate was cooled to room temperature on
 ice, 1 mg each of DNAse I and RNAse A were added, and
 the culture was allowed to stand at room temperature

for 30 minutes. Solid NaCl was added to 1 M and the

	current was praced on ice for I hour. Debris was
	removed from the lysate by centrifugation at 11,000
5	rpm in a Sorvall GSA rotor and polyethylene glycol
	(PEG) 8000 was added to a final concentration of 10 %
	(w/v). After 2 hours at 4°C, phage were collected by
	centrifugation as above and resuspended in a total
	volume of 15 ml of SM. The phage were extracted with
10	15 ml of CHCl3, centrifuged at 1600g for 15 minutes
	in an HB-4 rotor and the upper phase containing the
	phage was stored at 4°C overnight. Phage were
	purified by layering them on a step gradient
	consisting of 6 ml of 5 M CsCl in TM (10 mM Tris-HCl
15	pH 8.0, 10 mM MgCl2) layered over 6 ml of 3 M CsCl in
	TM. The gradient was centrifuged at 22,000 rpm in a
	Beckman SW28 rotor for 2 hours at 4°C. Phage banding
	at the 3 M/6 M CsCl interface were removed, mixed
	with an equal volume of saturated CsCl in TM and
20	layered in the bottom of an SW28 centrifuge tube.
	Phage were then sequentially overlaid with 3 ml of 6
	M CsCl in TM, 3 ml of 3 M CsCl in TM and sufficient
	TM to fill the centrifuge tube. The gradient was
	centrifuged as before and phage were recovered in the
25	same manner. Phage were dialyzed against three
	changes of 50 mM Tris-HCl, pH 8.0, 10 mM NaCl, 10 mM
	MgCl ₂ for one one hour each and then transferred to a
	polypropylene tube. The volume was adjusted to 1.2
	ml with dialysis buffer, and phage were lysed by
30	addition of 172 µl H ₂ O, 37.5 µl 20 % SDS, 60 µl 0.5 M
	Na_2EDTA , pH 8.0 and 30 μl of 5 $\mu g/m l$ proteinase K in
	water. After lysis for 1 hour at 55°C, phage DNA was
	extracted once with an equal volume of phenol, once
	with an equal volume of phenol:CHCl3 (1:1 v/v), and

once with an equal volume of CHCl3. DNA was precipitated by adding 80 µl of 3 M sodium acetate. pH 6.0 and 3.2 ml of ethanol and incubating the mixture for 5 minutes at room temperature. DNA was recovered by spooling it onto a pasteur pipet.

Spooled DNA was rinsed in 70 % ethanol and allowed to dissolve overnight by placing the pipet in 1 ml of 10 TE. pH 8.0.

Fragments of genomic clone 2-2 #4, were subcloned by partially digested 35 µg of 2-2 # 4 DNA with 80 units of Eco RI at 37°C. Time points of the

digestion containing 8.5 µg of DNA were removed at times ranging from 7.5 to 45 minutes of digestion and 15 Eco RI was inactivating by heating each time point to 70°C for 10 minutes. Small aliquots of time points were analyzed by electrophoresis in a 0.8 % agarose gel to determine the extent of digestion. Time

20 points showing partial Eco RI digestion products were ligated overnight with pUC18 DNA that had been cut to completion with Eco RI and dephosphorylated. Ligation reactions were diluted with 4 volumes of water and aliquots of each diluted reaction were used

to transform competent E. coli HB101. Aliquots of the transformation mixture were spread on LB plates containing ampicillin and plates were incubated overnight at 37°C. Plasmids from individual antibiotic resistant colonies were analyzed for 30 inserts containing Eco RI fragments of phage 2-2 #4 DNA. Large scale plasmid preparations were done from

subclones designated genomic 2-2 #2, 2-2#11, 2-2#17, and 2-2#23 whose inserts provide complete overlap of the region of the 2-2 #4 genomic clone that contained 35 the 2-2 gene (Figure 4A).

35 kbp fragment.

	The sequences of the cDNA clone In 2-2-3 and
	relevant portions of plasmid genomic subclone #2,
5	#11, #17 and #23 were determined by the method of
	Maxam and Gilbert (as described by Barker et al.,
	Plant Mol. Biol., 2, 335-350). The sequences of the
	genomic subclones were assembled to provide the
	complete nucleotide sequence of the 2-2 gene.
10	Comparison of the nucleotide sequences of the 2-2-3
	cDNA clone with the 2-2 #4 genomic sequence showed
	that 2-2 #4 contained a complete copy of the 2-2-3
	cDNA clone dispersed among several exons.
	The nucleotide sequence of the 5' untranslated
15	and promoter regions of the 2-2 #4 gene is shown in
	Figure 4B. The ATG functioning as the translation
	start codon for the 2-2 protein is contained within
	an natural Nco I site in the 2-2 #4 gene. Suitable
	promoter fragments weeful 6
20	promoter fragments useful for use in regulation the
	expression of recombinant DNA constructions can be
	removed from this subclone by cleavage of that Nco I
	site and removal of the promoter at any number of
	restriction sites 5' to that Nco I site such as at
25	Xho I to yield a 1.9 kbp fragment. Later examples
	teach the use of such fragments.
	A convenient Xho I site exists in genomic
	subclone 2-2 #11 nine nucleotides beyond the
	translation stop codon for the 2-2 protein (Figure
	4). Suitable downstream DNA fragments useful in
30	regulation the expression of chimeric genes can be
	removed from this subclone by cleavage of that Xho I
	site and removal of the downstream at any number of
	restriction sites 3' to that Xho I site such as Sal I
	to yield a 0.8 kbp fragment or Cla I to yield a 1.7

Example 3

5 Identification and Isolation of the Promoter and 3'
Downstream Regions of the 52.411 Corn 5-2 Gene

<u>Isolation and Characterization of 5-2 cDNA clones</u>

Details of the techniques used to perform

Example 3 are presented in Example 1. The cDNA

library made from poly (A)+ RNA from the roots of
N-(aminocarbony1)-2-chlorobenzenesulfonamide-treated
Missouri 17 corn plants described in Example 1 was
analyzed for additional cDNA clones representing
N-(aminocarbony1)-2-chlorobenzenesulfonamide-indused

N-(aminocarbony1)-2-chlorobenzenesulfonamide-induced mRNAs. The differential screening method described in Example 1 was used to isolate a new colony that displayed stronger hybridization with the cDNA probemade using RNA from roots treated with

20 N-(aminocarbonyl)-2-chlorobenzenesulfonamide. This colony was designated In 5-2.

A small scale plasmid preparation was performed on an overnight culture of In 5-2 and an aliquot of the plasmid, designated pIn 5-2, was nick-translated as described earlier. Slot blot analysis was performed as described in Example 1 using nick-translated In 5-2 plasmid. This analysis confirmed that pIn 5-2 contained a cDNA insert representing an mRNA that hybridizes strongly to RNA from N-(aminocarbonyl)-2-chlorobenzenesulfonamidetreated roots but not RNA from control roots. This

plasmid was designated pIn 5-2.

An aliquot of the small scale plasmid
preparation of pIn 5-2 was digested to completion
with Pst I and analysed by agarose gel
electrophoresis. The cDNA insert f the plasmid was

excised as a single 420 bp Pst I fragment. Plasmid pin 5-2 was nick-translated and used to probe a Northern blot of RNA from both untreated and N-(aminocarbony1)-2-chlorobenzenesulfonamide-treatedroots. The plasmid hybridized to a 2000 nt mRNA that was induced in root tissue by chemical treatment. As the insert of pln 5-2 was not a full length 10 copy of the 5-2 mRNA, the Agtll phage cDNA library made in Example 1 was screened for full length 5-2 cDNA clones. This was accomplished by probing the library with the purified cDNA insert from pIn 5-2 that had been nick-translated using the methods described in Example 1. Six different phage clones 15 showed homology to the pIn 5-2 cDNA insert and were plaque purified. Small scale DNA preps were made from these phage and aliquots of these DNAs were digested to completion with Eco RI and analyzed by 20 agarose gel electrophoresis. Three clones that contained insert similar in size to the 5-2 mRNA were subcloned into pUC18 by digestion of phage DNAs to completion with Eco RI and ligation of the resulting DNA into the Eco RI site of pUC18. One subclone, 25 designated pIn 5-2.32, was chosen for further analysis. Isolation of genomic clone 52.411 The library of Mol7 genomic DNA used to obtain 30 genomic clones for the 2-1 message was screened with nick-translated pIn 5-2 as described in Example 1 to isolate genomic clones corresponding to the 5-2 message. Six 5-2 genomic clones were plaque purified from this library in this manner. These genomic

clones were mapped by hybridization using a probe made from randomly primed cDNA synthesized using RNA

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from N-(aminocarbonyl)-2-chlorobenzene-

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sulfonamide treated corn roots to identify regions of homology to the In 5-2 cDNA as described in

Example 1. The results of this analysis indicated that all six clones appeared to contain the same regions of homology to the randomly primed cDNA probe. One clone, designated 52.411, was chosen for further analysis to determine its relationship to the In 5-2 cDNA.

Genomic clone 52.411 was digested to completion with Ecc RI and Sma I and the resulting fragments were ligated into the vector pUC 19 that had been cut to completion with the same two restriction endonucleases. Following transformation of E. coli with an aliquot of the ligation mixture, small scale plasmid preparations were performed on amp-resistant colonies that arose until a colony was found that

20 contained a 12 kbp Eco RI/Sma I fragment ligated into pUC 19. This plasmid was designated pJE 490.

The plasmid pJE 490 was digested to completion with Eco RI and Sal I and the resulting fragments were ligated into the vector pUC 19 that had been cut to completion with the same two restriction endonucleases. Following transformation of E. coli with an aliquot of the ligation mixture, small scale plasmid preparations were performed on amp-resistant colonies that arose until a colony was found that contained a 4 kbp Eco RI/Sal I fragment ligated into pUC 19. This plasmid, called pJE 491, Contains the 5' end of the 52.411 mess.

The plasmid pJE 490 was digested to completion with Sal I and the resulting fragments were ligated into the vector pUC 19 that had been cut to completion with the same restriction endonuclease. Following transformation of E. Coli with an aliquot

	92	
	of the ligation mixture, small scale plasmid	
5	preparations were performed on amp-resistant colonies that arose until one was found that contained a 4.0 kbp Sal I fragment ligated into pUC 19. This plasmid, called pJE 493, contains the 3' end of the 52.411 gene.	•
10	DNA Sequence Analysis of In 5-2.32 and 52.411 The sequence of the cDNA clone pIn 5-2.32 was	
15	method and Maxam and Gilbert chemical sequencing. Maxam and Gilbert chemical sequencing was performed on pIn 5-2.32 as described in earlier examples. For dideoxy sequencing, the plasmid pIn 5-2.32 as	
	separated by agarose gel electrophoronic and services	
20	cDNA insert was purified from the gel and dispested to completion with Sau 3A. The resulting DNA fragments were ligated into the New 1	
25	were ligated into the Bam HI site of the RF form of the vector MI3MP18. Aliquots of the transformation mixture were used to transfect E. coli JM 101. Aliquots of the transfection mixture were grown on 2XTT containing X-gal and IFTG. DNA was prepared from randomly chosen colorless plaques and sequenced by the dideoxy chain termination method using a	
30	Sequenase Kit® (U.S. Blochemicals) following the manufacturer's recommended protocols. The correct order of the Sau 3A fragments in pIn 5-2.32 was assigned by comparison of dideoxy sequence data from individual forms.	

Regions of the genomic DNA inserts contained 35 within the plasmids pJE 491 and pJE 493 were sequenced by creating nested sets of deletions of

by the Maxam and Gilbert method.

individual fragments with that derived for the cDNA

each plasmid as described in Example 1. By

- comparison of the sequences derived from regions of 5 pJE 491 to that derived from the In 5-2 cDNA clones, a 2.1 kbp Bam HI/Sal I genomic DNA fragment was identified containing 3.5 kbp of the 5-2 promoter as well as the start of the 5-2 structural gene (Figure 13). This fragment was subcloned into the vector 10 pBS(-). The resulting plasmid was designated pMC
- 3167.13. The sequence of the 5-2 gene upstream from the translation start of the 5-2 protein is shown in Figure 5.
- Site directed mutagenesis was performed on the 15 plasmid pMC 3167.13 to introduce a Nco I restriction site at the translation start of the 5-2 coding region. This was done so that the expression of a foreign coding sequence could easi<u>ly be placed under</u> the control of chemicals known to induce expression
- 20 of the 5-2 gene. An oligonucleotide of the sequence 5'-TGCCCATGGTGCGTG-3' was designed to introduce the Nco I site at the ATG codon corresponding to the start of the coding region of the 5-2 protein. The methods used to perform the mutagenesis were 25
- described in Example 1. The resulting plasmid Containing the mutagenized 5-2 promoter was designated pMC 75.j5, and is shown in Figure 6.

EXAMPLE 4

30 Identification, Isolation and Modification of corn 218 gene Promoter

Isolation and Characterization of 218 cDNA clones

Details of the techniques used to perform the 35 procedures used in this Example are presented in Example 1. Equimolar aliquots of the cDNA used to

	make the Agtll phage cDNA library in Example 1 and	
	pucis DNA that had been digested to completion with	
5	ECO RI and dephosphorylated were ligated together	
	overnight. Aliquots of the ligation mixture were	,
	transformed into competent E. coli DH5 cells (RDL)	•
	and plated onto LB plates containing 50 mg/ml	
	ampicillin. Antibiotic resistant colonies were	
10	arrayed onto nitrocellulose disks and analyzed for	
	cDNA clones containing inserts representing mRNAs	
	induced by substituted benzenesulfonamides as	
	described in Example 1. A colony displaying stronger	
	hybridization with the cDNA probe made from RNA of	
15	N-(aminocarbonyl)-2-chlorobenzenesulfonamide-treated	
	Corn roots was identified. This clone was designated	
	In 218 and the plasmid contained within it was	
	designated p218. Agarose gel electrophoresis of the	
	Eco RI digestion products of p218 showed the plasmid	
20	Contained a 900 bp insert. Hybridization of	
	nick-translated p218 to size fractionated RNA	
	isolated from N-(aminocarbonyl)-2-chloro-	
	benzenesulfonamide-treated roots indicated that the	
	cDNA was full length.	
25	A library of Missouri 17 genomic DNA was made	
	and screened for genomic sequences corresponding to	
	the 218 cDNA clone using nick-translated p218 as	
	described in Promple 1 and 1	
	described in Example 1 with the following changes: 1)	
30	genomic DNA was digested with Eco RI rather than Sau	
	3A and 2) Eco RI fragments of the appropriate size	
	were cloned in the vector \Dash that had been	
	digested with Eco RI rather than using Bam HI	
	general denomic clones unbridizing to the	
35	218 cDNA were identified and plaque purified from	,
	this library. The Eco RI inserts from members of	•
	each group were subcloned into the plasmid vector	

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pBS(+) and the subcloned genomic DNA was digested with a variety of restriction ensymes. The digestion products were separated by agarose gel electrophoresis, blotted to nitrocellulose and probed with nick-translated pIn218. Comparison of the restriction maps generated for the genomic subclones with that derived for the 218 cDNA indicated that one genomic subclone, designated pMC730, contained a 1.4 kbp Sac I/Xho I fragment that was very similar to and hybridized that hybridized with the 218 cDNA clone.

Plasmid pMC730 was digested to completion with Xho I and the the reaction mixture was diluted to 200 ul. After heating at 65°C to inactivate XhoI, the diluted digest was ligated together to recircularize the plamid, and thus deleting a 6 kbp Xho I fragment

This plasmid was designated pMC767. The plasmid
pMC767 clone was sequenced 224 bases from the XhoI
side and was found to compare well with the cDNA for
190 bases at which point an intron junction was
encountered. To skip over this intron, pMC767 was
digested to completion with Nco I and Xho I. The 5'
overhanging ends were rendered blunt using I4
polymerase and the plasmid was recipally of

from pMC730 that did not hydridize with the cDNA.

polymerase and the plasmid was recircularized as described above to create the plasmid pMC791 (Figure 7).

From this plasmid a DNase I deletion series was generated for dideoxy sequencing. The entire Nco I to Eco RI region was sequenced (1710 bases) and compared with the cDNA (Figure 8). The genomic sequence matched the cDNA sequence at its 5 end and extended over 1.5 kb beyond the 5 end of the cDNA (Figure 7). The beginning of the 218 message was determined by using the genomic clone in a riboprobe

protection experiment and the first ATG of the message was identified by searching downstream from 5 this site, and is indicated at nucleotide 1516 by an arrow in Figure 7. Computer analysis of the genomic sequence identified an Afl III site that contained this ATG (underlined in Figure 7). Digestion with this enzyme produces a cohesive end containing the 10 ATG start codon of the 218 gene product that is capable of ligating with any desired coding region. Thus a functional 1.4 kbp 218 promoter and 5 untranslated leader fragment may be obtained from pMC791 by partial Afl III digestion followed by 15 complete digestion with SmaI to excise a 1.4 kbp

EXAMPLE 5

20 Identification, Isolation, and Modification of the Promoter and 3' Downstream Regions of the P6.1
Petunia Gene

Growth and Chemical Treatment of Plants

promoter/untranslated leader fragment.

25 Petunia (Mitchell) seeds were germinated in soil and allowed to grow for one month under standard greenhouse conditions. Plants were transferred to a hydroponic growth apparatus in a greenhouse using foam plugs to support the plants. These plugs were foam placed in holes in a wooden bpard and placed over a stainless steel sink containing 0.5X Hosgland's solution. The solution was aerated using standard aquarium pumps, and was changed weekly.

After one month of hydroponic growth, plants were

35 transferred to stainless steel trays containing either fresh 0.5 X Hoagland's or 0.5X Hoagland's

containing 0.2 g per liter of N-(aminocarbonyl)-2-chlorobenzenesulfonamide. Root tissue was

harvested after six hours of treatment.

Isolation of RNA

Root tissue was harvested by slicing roots off just below the foam plugs. Tissue (2-5 g) was 10 wrapped in aluminum foil, quick frozen in liquid nitrogen and stored at -80°C until used. Prozen tissue was transferred to a mortar pre-cooled with liquid N2 and ground to a fine powder with a chilled pestle. The powder was transferred to a 50 ml polyethylene centrifuge tubes containing 10 ml NTES (0.01 M Tris-HCl pH 7.5, 0.1 M NaCl, 1 mM EDTA.1% SDS). 10 ml waters was transferred.

SDS), 10 ml water-saturated phenol, and 10 ml chloroform: isoamyl alcohol (24:1 v/v). The emulsion was vigorously shaken for 15-30 minutes and then

- separated by centrifugation in 30 ml lorex® tubes at 5000 rpm for 10 minutes in a Sorvall HB-4 rotor.

 Nucleic acids were precipitated from the aqueous phase by the addition of 1 ml 3M sodium acetate, pH 6.0 and 25 ml ethanol. After 2 hours at -20°C, the precipitate was collected by centrifugation at 10,000
- rpm for 20 minutes in a Sorvell S834 rotor. Pellets were drained well and dissolved in 2 ml of H₂O. Two ml of 4 M lithium acetate was added to selectively precipitate the RNA and the solution was held on ice for 3 hours. RNA was collected by centrifugation at
- 10,000 rpm in an SS-34 rotor for 20 minutes

 RNA was dissolved in 400 µl water, transferred to 1.5

 ml microcentrifuge tubes and reprecipitated with

 ethanol for 2 hours at -20°C. RNA was collected by
- 35 centrifugation in a microcentrifuge for 5 minutes and the final pellets were dissolved in 200 µl H₂O. RNA

concentrations were determined from the absorbance of the solutions at 260 nm. Yields of RNA from typical preparations were approximately 1 mg.

Isolation of Poly(A)+ RNA

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Poly(A)+ RNA was purified from total RNA by oligo (dT) cellulose chromatography. 2.5 mg of RNA 10 was diluted to 0.4 mg/ml (10 A260 per ml) in zero salt buffer (10 mM Tris-HCl pH 7.4, 0.5% SDS, 1 mM EDTA). The RNA was denatured at 65°C for 5 minutes and then chilled on ice for 10 minutes. Sodium chloride was then added to bring the concentration to 15 0.4 M. The RNA was applied to a plastic disposable

- column that was packed with 0.1 g oligo (dT) cellulose (Worthington) which had been equilibrated with high salt buffer (zero salt buffer containing 0.4 M sodium chloride). RNA was passed over the column two or
- three times to maximize binding of the poly(A)+ fraction. Following binding, the column was washed with 10 ml high salt buffer. Poly(A)+ was eluted with zero salt buffer in 6 one ml fractions. Absorbance of the fractions was measured at 260 nm
- 25 and the fractions containing RNA were pooled. RNA was precipitated with ethanol and dissolved in 100 ul H2O. Yields of poly(A) + RNA were generally 0.5 - 1% of the total RNA applied to the column.

30 Construction of cDNA Library

Five ug of poly(A) + RNA from N-(aminocarbonv1)-2-chlorobenzenesulfonamide-treated roots were ethanol precipitated, collected by centrifugation and dissolved in 10 µl of H2O. The

35 RNA was heated at 65°C for 3 minut s and rapidly chilled on ice. First strand cDNA was prepared using

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a reaction mixture containing 10 μl RNA, 5 μl 10 χ first strand buffer (0.5 M Tris-HCl pH 8.5, 0.4 M RCl, 0.1 M MgCl2, 4 mM DTT), 5 µl of a nucleotide mixture containing each of the four dNTPs(ACGT) at 10 mM, 5 μ l 100 μ g/ml oligo (dT)₁₂₋₁₈, 5 μ l α -³²P dCTP, 2 µl placental rbonuclease inhibitor and 50 units of reverse transcriptase. The reaction was incubated at 42°C for 1 hour. The mass of cDNA synthesized was calculated from the incorporation of $^{32}P-dCTP$ into the synthesized DNA. The RNA:cDNA duplex was denatured by heating in a boiling water bath for 1.5 minutes, then quick chilled on ice. The following were then added to the 50 μ l first strand reaction mixture: 50 µl 2X second strand buffer (100 mM HEPES pH 6.9, 100 mM KC1, 20 mM MgCl $_2$), 1 μ l of a 10 mM dNTP mixture and 2 μl DNA polymerase 1 (50 $U/\mu l$). The reaction mix was incubated at 15°C for 5 hours. At that time, 400 μl of S1 buffer (30 mM sodium acetate pH 4.4, 250 mM sodium chloride, 1 mM ZnCl2) and 500 units of Sl nuclease were added. The incubation was continued for 1 hour at 37°C. The products of the S1 reaction were extracted with an equal volume of phenol:chloroform (1:1 v/v) and precipitated with etharol. The pellet was dissolved in 20 µl methylase buffer (50 mM Tris-HCl pH 7.5, 1 mM EDTA, 5 mM DTT) to which 2 µl 100 mM S-adenosylmethionine and 1 μl of Eco RI methylase (40 U/µl) were added. The methylation reaction was incubated at 37°C for 15 minutes followed by 65°C for 10 minutes. The ends of the cDNA were filled in by adding 2.5 µ1 0.1 M MgCl2, 2.5 µ1 0.2 mM d(ACGT)TP and 1 μ 1 DNA polymerase 1 (5U/ μ 1) to the tube and allowing the fill-in reaction to proceed for 20 minutes at room temperature. The cDNA was then

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extracted with phenol:chloroform (1:1 v/v) and ethanol precipitated. The pellet was dissolved in 32 5 μ1 H₂O, 10 μ1 phosphorylated Eco RI linkers (0.1 mg/ml), 5 μ l 10% ligase buffer, and 3 μ l of T4 DNA ligase (0.1 ml) (6 Weiss units/ μ l). The ligation reaction was then incubated at 15°C for 16 hours. The DNA ligase was inactivated by heating at 65°C for 10 10 minutes and Eco RI linkers were digested for 2 hours at 37°C by adding 40 µl H2O, 10 µl 10% Eco RI buffer and 3 µl Eco RI (20 U/µl) to the DNA. The cDNA was then precipitated with ethanol, dissolved in 20 µl 1X TBE and subjected to electrophoresis in a 6 15 % polyacrylamide gel. The gel was stained with ethidium bromide (1 μ g/ml) to visualize the cDNA in the gel. A slice of the gel containing cDNA >0.5 kbp was cut out and DNA was recovered by electroelution of the cDNA into a dialysis bag. The electroeluted cDNA was extracted with phenol:chloroform (1:1 v/v), 20 precipitated with ethanol, and dissolved in 20 µ1 HoO. One µl of the cDNA was counted in a liquid scintillation spectrometer and the mass of cDNA was determined using the specific radioactivity of the 25 32p-dCTP used in the cDNA synthesis. One microgram of Agt10 arms that had been cut to completion with Eco RI and dephosphorylated was ligated to 30 ng of cDNA in a volume of 5 μ l. The ligation mixture was then packaged using Gigapack extracts (Stratagene) as

Isolation of cDNA clone P6

procedure.

Approximately 10,000 phage were plated ut on 5 150 mM LB agar plates containing 10 mM MgCl2 (2000

per manufacturer's instructions. Approximately 1 million recombinants were obtained from such a

		phage per plate) using the E.coli strain C600 as the
		host. Replica filters copies of the library were
	5	prepared from each plate as follows: Dry
•		nitrocellulose filters were wetted by placing them
:		onto the surfaces of agar plates containing the phage
		cDNA library. The filters were then transferred to
		a sheet of Whatman 3MM paper that had been saturated
	10	with 0.5 M NaOH and 1.5 M NaCl for 30 seconds to 1
		minute. The filters were transferred to a sheet of
		Whatman 3MM that had been saturated with 1 M Tris-HC1
		pH 7.0 and 1.5 M NaCl for 5 minutes, rinsed in 2X
		SSC, air dried for 1 hour and baked in vacuo for 2
	15	hours at 80°C. This process was repeated for each
		plate to make multiple filter copies of the library.
		The replica filters of the cDNA library were
		screened for cDNA clones representing mRNAs induced
		by N-(aminocarbony1)-2-chlorobenzenesulfonamide by
	20	the differential hybridization method described in
		Example 1. cDNA probes were prepared from poly(A)+
		RNA from both untreated and treated root tissue as
		described for first-strand cDNA synthesis in this
		example with the following modifications: One
	25	microgram of poly(A)+ RNA, 2.5 µl of 1 mM dCTP
		and 10 µl 32p-dCTP (10 mCi/ml) were used in the
		reaction. Following probe synthesis, the RNA
		template was hydrolyzed by the addition of 25 µ1 0.15
		M NaOH and incubating the cDNA at 65°C for 1 hour.
	30	Base was neutralized by addition of 12.5 µl 2 M
		Tris-HCl pH 8.0 and 25 µl l N HCl. Single-stranded
		cDNA was separated from unincorporated label on a
•		Sephadex® G50 column, equilibrated and run in 10 mM
:		Tris-HCl pH 7.5, 1 mM EDTA. Fractions eluting in the
•	35	void volume were pooled, thanol precipitated and

dissolved in H2O.

Replica filters were prehybridized in a solution of 0.1% SDS, 4X SSC, 5X Denhardt's solution, 50 mM sodium phosphate pH 6.8 at 42°C for 5 hours.

- 5 50 mM sodium phosphate pH 6.8 at 42°C for 5 hours. The solution was replaced with hybridization buffer (prehybridization buffer containing 50% deionized formamide) containing 5 x 10° cpm/ml of probe using RNA from either untreated or N-(aminocarbonyl)-2-
- 10 chlorobenzenesulfonamide-treated roots.

 Hybridizations were incubated for 24 hours at 42°C.

 The filters were then washed twice at room
 temperature for 1 hour with 2x SSC, 0.1% SDS. A
 final wash was conducted at 50°C in 0.1% SSC, 0.1%
- SDS for one additional hour. Filters were exposed to X-ray film at -80°C for 60 hours with one intensifying screen.

Plaques hybridizing more strongly with the probe derived from N-(sminocarbonyl)-2-chloro-

- 20 benzenesulfonamide-treated roots were deemed positive clones in the differential screen. These plaques were removed from the plates with 100 μl capillary pipets and placed in 0.5 ml of SM. Plaque purification was performed on these phage as
- 25 described in Example 1 by repeated differential screening using the hybridization procedure described above. One clone purified in this manner was designated P6.
- A liquid lysate of P6 phage was prepared by
 30 absorbing 10% of the phage eluted from one plaque to
 100 µl of an overnight culture of E.coli BNN102, and
 inoculating 30 ml NZCYM (per liter: 10 g NZ amine, 5
 g yeast extract, 5 g NaCl, 1 g cosamino acids, 2 g
 MySO₄, pH 7.5) with the resulting infected culture.
 35
- 35 After 5 hours of growth at 37°C, complete lysis of the bacteria had occurred. The lysate was cleared by

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centrifugation at 10,000 rpm for 10 minutes in a Sorvall SS34 rotor, and the supernatant was

- 5 transferred to a clean tube. RNAse A and DNAse I were added to 10 μg/ml and 20 μg/ml respectively and the lysate was incubated for 15 minutes at 37°C. One-fifth volume of 20% PEG 6000, 2.5 M NaCl were added to the lysate and phage were allowed to properly table for 15 minutes.
- 10 precipitate for 15 minutes at room temperature. The phage were collected at 10,000 rpm for 10 minutes, and the pellet was drained well. Phage were resuspended in 0.5 ml 45 PEG 6000, 0.5% NaCl and transferred to a microfuge tube. The phage were
- extracted with 0.5 ml phenol:chloroform (1:1 v/v) and DNA was precipitated with 2 volumes of ethanol. DNA was collected by centrifugation and dissolved in 50 µl TE pH 8.0. Five µl of DNA were digested to
- completion with Eco R1 and resulting DNA fragments 20 were analyzed by agarose gel electrophoresis. The results of this analysis showed that the P6 cDNA clone contained a single 700 bp insert.

The Eco R1 insert of P6 was subcloned from the phage vector Agt10 to the plasmid pUC119. Ten µg of P6 DNA was digested to completion with Eco R1 and digestion products were subjected to electrophoresis on a 1% agarose gel. A piece of the gel containing the 700 bp Eco R1 fragment was cut out and placed in a piece of dislysis tubing containing 0.5 ml 1X TAE (0.04 M Tris-HC1 pH 7.8, 2 mM EDTA). The DNA was electrocluted from the gel piece at 100 volts for 15 minutes. The buffer containing the DNA was removed

from the bag, extracted with an equal volume of phenol:chloroform (1:1 v/v), and DNA was precipitated with ethanol in the presence of 0.3 M sodium acetate. Ten µg of pUC:19 was digested to completion

with Eco Rl. extracted with phenol:chloroform (1:1 v/v), and precipitated with ethanol. Equimolar 5 amounts of vector and insert were ligated in a volume of 10 µl at 15°C for 2 hours. An aliquot of the ligation mixture was used to transform competent E. coli JM83 cells. Aliquots of the transformation mixture were grown overnight at 37°C on LB plates 10 containing 75 µg/ml ampicillin that had been spread with X-Gal and IPTG. Small scale plamid preparations were performed on white colonies and aliquots of the DNAs were digested to completion with Eco Rl until one was found containing the desired 700 bp Eco RI 15 fragment from P6 in pUC 119. The resulting clone was designated P6.1. Ten μ g (2 mg/ml) of total RNA from control and N-(sminocarbonyl)-2-chlorobenzenesulfonamide-treated roots was denatured by adding 10 µl of deionized 20 formamide, 3.5 µl formaldehyde, 4 µl 5X MEN buffer (40 mM MOPS pH 7.0, 10 mM sodium acetate, 1 mM EDTA) and incubating at 65°C for 15 minutes. The RNA was subjected to electrophoresis in a 1.5% agarose gel containing formaldehyde and 1X MEN until the 25 bromphenol blue had migrated to the bottom of the gel. RNA was stained in the presence of 10 mM sodium phosphate pH 6.8 and 1 µg/ml acridine orange for 30 minutes. The gel was then destained in 10 mM sodium phosphate for 30 minutes, and the RNA was visualized 30 on a UV transilluminator, photographed, and blotted to nitrocellulose (Millipore HAWP). To do this. Whatman 3MM paper was placed below the gel on a glass plate so that the ends of the paper extended into 20% SSC. A sheet f nitrocellulose which had been prewet 35 with 2X SSC was placed on top of the gel followed by a layer of Whatman 3MM, then a stack of paper towels

		10 cm high. A glass plate and weight were then
		placed on top of the stack. Following an overnight
	5	transfer, the filter was rinsed briefly in 2% SSC,
:		air dried, and baked in vacuo for 2 hours at 80° C.
_		The filters were prehybridized for 5 hours in
4		plastic dishes at 42°C using the hybridization buffer
		described earlier. Plasmid p6.1 was nick-translated
	10	by combining 1 µg of DNA with 5 µl 10X buffer (0.5 M
		Tris-HCl pH 8.0, 0.1 M MgSO ₄ , 10 mM DTT and 0.5 mg/ml
		BSA), 5 µl 0.3 uM d(AGT)TP, 5 µl ³² P-dCTP (Amersham,
		10 mCi/ml 400 Ci/mm-l-) a p-dCTP (Amersham,
		10 mCi/ml, 400 Ci/mmole), 1 µ1 DNA polymerase 1 (5
	15	U/μl, Boehringer-Mannheim), and 1 μl of 0.1 μg/ml
		DNAse 1 in a total volume of 50 µl. The mixture was
		incubated for 1.5 hours at 14°C, and the reaction was
		stopped by the addition of 5 μl of 0.25 M EDTA. The
		reaction was then incubated for 5 minutes at 70°C,
		and labelled DNA was separated from unincorporated
	20	nucleotides by Sephadex® G-50 column chromatography.
		The prehybridization solution was removed from the
		bag and replaced with hybridization solution
		containing nick-translated plasmid P6.1 DNA at a
		concentration of 1 x 106 cpm/ml. Hybridization was
	25	carried out at 42°C for 24 hours on shaking
		platform. Filters were washed twice with 2X SSC,
		0.1% SDS at 42°C followed by two washes in 0.1% SSC,
		0.1%SDS at 60°C. The filter was wrapped in
		polyethylene food wrap and exposed to X-ray film at
	30	-80° C for 16 hours with one intensifying screen .
		The P6.1 probe hybridized to an 800 bp message
		in RNA from N-(aminocarbonly)-2-chlorobenzene-
;		sulfonamide-treated roots while no signal was
		observed in RNA from untreated plants. The insert
k.		prants. The Insert

35 size of the cDNA clone approximated the size f the

hybridizing RNA, indicating that P6.1 was potentially

a full-length cDNA clone. 5 Sequence Analysis of the cDNA clone P6.1 The nucleotide sequence of clone P6 was determined by sequencing a nested set of deletions mutants generated by digestion of the cDNA insert in 10 P6.1 with Exo III nuclease. The Eco RI insert from cDNA clone P6.1 was subcloned into the Eco RI site of the vector Bluescript(-) (Stratagene). The resulting clone was designated P612. Ten µg of P612 DNA were digested with Kpn I (3' overhang which is resistant 15 to Exc III digestion) and Kho I (5' overhang which is sensitive to Exo III). The DNA was extracted with phenol:chloroform (1:1 v/v), precipitated with ethanol then resuspended in 63.5 ul of H-O. Eight ul of 10X Exo III buffer (0.5 M Tris-HCl pH 8.0, 50 mM 20 MgCl₂, 100 mM β -mercaptoethanol) and 3 μ l of Exo III (100 $U/\mu l$) were added and the mixture was incubated at 37°C. Aliquots of 2.5 µl were removed every 30 seconds for 15 minutes and added to 13.5 µl ice cold quenching buffer (100 mM sodium acetate pH 4.7, 600 mM NaCl, 20 mM zinc acetate). The aliquots were 25 pooled into groups of five sequential time points and treated with 1 unit of SI nuclease at room temperature for 30 minutes. Water (123 µl) was added to each pool, and 10 µl from each was analyzed by 30 agarose gel electrophoresis. The remaining DNA was extracted with phenol:chloroform (1:1 v/v), precipitated with ethanol and resuspended in 20 ul of fill-in/ligation buffer (20 mM Tris-HCl pH 7.8, 25 mM

NaCl, 10 mM MgCl2, 20 mM DTT, 1 mM ATP, 0.1 mM

dNTPs). Forty units of DNA ligase and 2 units of Klenow fragment were added and the mixture was

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	incubated overnight at 15°C. Ten µl of the ligation
	mixture was used to transform competent E. coli hos
. 5	MV1193 cell and aliquots of the transformation
;	mixture were spread onto LB plates containing 75
	µg/ml amp. Ten colonies from each transformation
:	were analyzed for insert size, and a series of clone
	were selected for sequencing that represented
10	deletions of the initial cDNA insert that were each
	Drogressively lee harries con insert that were each
	progressively 150 base pairs longer. Single-strande
	DNA from those clones were sequenced using the M13
	reverse primer and the method of dideoxy chain
15	termination described in Example 1.
13	*
	Isolation of a genomic clone corresponding to P6 cDN
	Twenty grams of petunia leaf material was
	harvested, submerged in ice water and transferred to
	a Chilled mortar. Twenty ml of Buffer A (10 mM
20	Tricine pH 7.6, 1.4 M sucrose, 5 mM MgCla, 5 mM
	β-mercaptoethanol) was added to the mortar and leaf
	tissue was ground to a fine pulp. The solution was
	diluted to 100 ml with Buffer A and filtered through
	four layers of cheesecloth. The filtrate was then
25	passed through eight layers of cheesecloth and
	centrifuged at 2500 rpm for 10 minutes in a Sorvall
	GSA rotor. The pellet was resuspended in 100 ml
	Buffer A and centrifuged as before. The pellet was
	resuspended in 100 ml Buffer B (Buffer A containing
30	0.4% Triton X-100), held at 4°C for 10 minutes,
	centrifuged as before. The resulting pellet was
	resuspended in 100 ml of Buffer B and the
:	centrifugation was repeated at 2000 rpm for 10
	minutes wielding at 2000 rpm for 10
*	minutes, yielding a crude nuclear pellet. This

pellet was resuspended in 4 ml 50 mM Tris-HCl pH 8.0

and 20 mM EDTA to which 0.5 ml of 10% sarkosyl was

	added. The solution was incubated at 60°C for 5
	minutes, and then cooled to room temperature.
5	One-tenth ml of a 5 mg/ml proteinase K solution was
	added and the incubation was continued at 37°C for 4
	hours with gentle shaking. The volume of the
	solution was measured and 1 g solid cesium chloride
	was added per 1.2 ml of solution. Ethidium bromide
10	was added to 0.5 mg/ml and the density adjusted to
	1.55 g/ml with CsCl. The DNA was banded by
	centrifugation at 40,000 rpm for 30 hr at 15°C in a
	Beckman 70.1Ti rotor. The band was collected from
	the CsCl gradient by side puncturing of the
15	centrifuge tube. Ethidium bromide was removed from
	the DNA by repeated extraction with isoamyl alcohol
	equilibrated with TE pH 8.0. The DNA was then
	dialyzed against 5 mM Tris-HCl pH 8.0, 0.25 mm EDTA
	for 2 days.
20	Conditions were established for partial
	digestion of petunia genomic DNA by performing pilot
	restriction digests. Ten µg of DNA was brought up to
	a volume of 150 µl with the appropriate restriction
	buffer. Thirty µl aliquots of the DNA was dispensed
25	into a microcentrifuge tube labelled #1. Fifteen µl
	were dispensed into seven tubes labelled #2-8, and
	the remainder into tube #9. All tubes were chilled
	on ice. Sau 3A (4 units) was added to tube #1 and
	the contents of the tube were mixed well. Fifteen µl
30	from tube #1 was added to tube #2. This twofold
	serial dilution was continued through to tube #8, and
	all tubes incubated at 37°C for 1 hour. The
	restriction digestions were stopped by chilling the
	tubes to 0° C and adding EDTA to 20 mM. The samples
35	were subjected to electrophoresis through a cos

agarose gel at 1-2 V/cm. The enzyme concentration

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which yielded maximum intensity of fluorescence in the 15-20 kbp range was determined after ethidium bromide staining of the gel. Half of the enzyme/DNA ratio determined above was chosen for the preparative digestion of genomic DNA in order to maximize yield of DNA fragments in the 15-20 kbp size range. That enzyme concentration ranged between 0.06 and 0.25 10 units of Sau 3A per ug DNA. Three hundred µg of DNA was divided into 3 tubes: 1/4 in tube #1, 1/2 in tube #2 and 1/4 in tube #3, and the concentration of DNA was adjusted to 67 ug/ml. Sau 3A was added to tube #2 at the final 15 concentration which was thought to maximize for 15-20 kb molecules. Tube #1 contained one half that concentration while tube #3 contained twice as much All reactions were incubated at 37°C for 1 hour. After stopping the digestion as above, aliquots 20 from each of the digestions were analyzed by agarose gel electrophoresis and the appropriate digestions containing maximum amounts of 15-20 kbp fragments were pooled. The pooled sample was loaded onto a 10-40% sucrose gradient in 1 M NaCl, 20 mM Tris-HCl 25 pH 8 and 5 mM EDTA and centrifuged at 26,000 rpm for 24 hours at 20°C in an Beckman SW41 rotor. Fractions of 0.5 ml were collected from the gradient and 15 μl of every third fraction were analyzed by agarose gel electrophoresis. Fractions containing 15-20 kbp DNA 30 fragments were pooled and dialyzed against 4 liters TE for 16 hours at 4°C. After dialysis, the volume of DNA was reduced to 3-5 ml by repeated extraction with 2-butanol, followed by precipitation of the DNA with ethanol in the presence of 0.3 M sodium 35 acetate. The DNA was dissolved in TE at a

concentration of 300-500 µg/ml.

	Genomic DNA was ligated to Bam HI cut and	
	dephosphorylated EMBL3 arms (Stratagene) according to	
5	the manufacturer's instructions using 2 fold molar	,
	excess of vector to insert. The ligation was	6
	packaged using Gigapack extracts (Stratagene). A	
	library was plated by adsorbing 20,000 phage to 350	•
	μl of an overnight culture of E. coli LE392 for 15	
10	minutes at 37°C. A 7.5 ml aliquot of molten top	
	agarose (LB plus 0.8% agarose at 50°C) was added the	
	bacteria and the culture was spread on 150 mm LB	
	plates containing 10 mM MgSO4. A total library of	
	260,000 phage was plated in this manner.	
15	The genomic library was screened for P6 genomic	
	clones using the cDNA insert from the P6.1 clone as a	
	probe. To do this, the insert was cloned into the	
	transcription vector BS(-) (Stratagene). Ten µg of	
	P6.1 was digested to completion with Eco R1 and the	
20	resulting DNA fragments were separated by agarose gel	
	electrophoresis. The cDNA insert fragment was	
	electroeluted from the gel, extracted with an equal	
	volume of phenol:chloroform (1:1 v/v) and	
	precipitated with ethanol. Ten µg of vector pBS(-)	
25	DNA was digested to completion with Eco RI, with	
	extracted phenol:chloroform (1:1 v/v) and	
	precipitated with ethanol. Insert and vector were	
	ligated together in a final volume of 10 µl for 2	
	hours at 15°C and an aliquot of the ligation mixture	
30	was then used to transform competent E. coli JM83.	
	The transformation mixture was plated out on LB	
	plates containing 75 g/ml ampicillin which had been	•
	spread with X-gal and IPTG prior to plating of	•
	bacteria. Small scale plasmid preparations were	:
35	performed on white colonies and DNAs were digested	•
	with Eco R1. A colony containing the desired P6 cDNA	

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insert in the vector pBS(-) was identified and named

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- P6.11 was linearized by digestion with Bam HI and α -32p UTP labelled RNA transcript was made from the plasmid using T3 polymerase following the manufacturer's protocols (Fromeoa Biotech Inc.).
- Nitrocellulose replicas of the petunia genomic

 10 library were made and prehybridized for 3 hours as
 described earlier. The prehybridization solution was
 replaced with hybridization solution containing the
 p6.11 RNA probe at 2 x 10⁶ cpm/m1. Hybridization was
- performed for 24 hours at 42°C with gentle
 15 agitation. The filters were washed twice with 2x
 SSC, 0.1% SDS at 42°C, followed by two washes with
 0.1% SSC, 0.1% SDS at 42°C. The filters were exposed
 to X-rey film at -80°C for 24 hours using a single
- intensifying screen. Three phage displayed strong
 phybridization to the probe were plaque purified as
 described earlier and designated phage 1, 2 and 3.

Characterization of genomic clones

Phage were grown in liquid culture by
inoculating 300 ml of NZCYM media with 10¹⁰ phage
which had been previously adsorbed onto 1 ml of an
overnight culture of E. coli LE392. The infected
culture was grown at 37°C with shaking until complete
lysis of bacteria occurred (generally by 7 hours).

- 30 Cellular debris was removed from the lysate by centrifugation, and the supernatant treated with 1 μg/ml of both DNAse and RNAse for 1 hour at room temperature. Solid sodium chloride was added to 1 M, and PEG 6000 added to 10% (ω/ν). The phage were
- 35 allowed to precipitate overnight at 4°C, and then collected by centrifugation in a Sorvall GSA rotor at

7000 rpm for 15 minutes. The phage pellets were resuspended in SM and 0.75 gram cesium chloride was added per ml SM. Gradients were centrifuged in a Beckman 70.1Ti rotor at 38,000 rpm for 24 hours at Phage bands were collected from the sides of the tubes and dialyzed overnight at 4°C against 10 mM NaCl, 50 mM Tris-HCl pH 8, 10 mM MgCl2. DNA was 10 extracted from purified phage by adding sodium chloride to 20 mM, promase to 0.5 mg/ml and SDS to 0.5% followed by incubation of the resulting solution at 37°C for 1 hour. The sample was dialyzed against TE pH 8 and precipitated with ethanol. This yielded 15 approximately 250 ug of phage DNA. Phage DNAs were digested with Sal I to excise the insert DNA from vector. Agarose gel electrophoresis of <u>digested DNA showed that phage 1,</u> 2 and 3 contained inserts of 13, 14 and 10 kb 20 respectively. By further restriction enzyme digestions and hybridizations to the cDNA clone P6.1. restriction maps were generated that indicated that the inserts of all three phage overlapped one another and were fragments of the same region of petunia DNA. 25 Two Eco R1 fragments of 0.6 kbp and 1.8 kbp from phage 1 and phage 3 were found to hybridize to cDNA P6.1 in mapping experiments described above. These fragments were separated by a 5 kbp nonhybridizing Eco RI fragments. This suggested either the presence of a large intron in the P6 gene or the 30 existence of two genes homologous P6.1 on the same genomic DNA fragment. To address these possibilities, the two Eco Rl fragments were subcloned and sequenced. T n µg of phage 1 DNA were 35 digested to completion with Eco Rl and the products

were separated by agarose gel electrophoresis. The

- 1.8 and 0.6 kbp fragments were electroeluted from the gel, extracted with phenol:chloroform (1:1 v/v) and 5 precipitated with ethanol. Each fragment was ligated into Eco RI digested pUC119 DNA in a final volumes of 10 ul for 2 hours at 15°C. The ligations mixtures were used to transform competent E. coli JM83 cells. Aliquots of the transformation mixture were plated 10 out on LB plates containing 75 g/ml ampicillin which had been spread with X-gal and IPTG prior to plating of bacteria. Small scale plasmid preparations were performed on white colonies and the resulting DNAs were digested with Eco Rl. Subclones containing the 15 desired 0.6 and 1.8 kbp fragments were chosen in both possible orientations to facilitate sequencing the ends of the fragments. These two orientations were identified by digesting subclones with Sal I for the
- 0.6 kbp fragment and Pvu II for 1.8 kbp fragment.

 The resulting plasmids were designated P619 and P620 (two orientations of the 0.6 kbp genomic fragment) and P621 and P622 (two orientations of the 1.8 kb genomic fragment).
- Plasmid DNAs were sequenced by dideoxy chain

 25 termination method using 35s-dATP as described in
 earlier examples. Sequence analyses showed that the
 1.8 kbp Eco RI genomic fragment contained a gene with
 perfect homology to the Pf.1 cDNA while the 0.6 kbp
 genomic fragment contained a closely related gene.

 30 The homologous gene is the 1.8 kbp EcoR I fragment

Mapping the endpoints of Gene 1

was designated gene P6.1.

A primer extension analysis was performed to

determine the 5' end of the P6 RNA. An
oligonucleotide complementary to the coding strand in

	the P6 gene from 12-33 bases downstream of the first	
	in-frame ATG was synthesized using an Applied	
5	Biosystems DNA Synthesizer. The oligonucleotide,	
	5'-CCACTAAGACAATCTAAAGACC-3' was end-labelled with	
	32 P by drying 50 uCi of α - 32 ATP in a microfuge tube	
	using a Speedvac centrifuge. Two µl of	•
	olignucleotide (2.5 pmole/µ1), 2 µ1 5X kinase buffer	
10	(125 mM Tris-HC1 pH 9.5, 25 mM MgCl2, 12.5 mM DTT,	
	2.5 mM spermidine, 0.25 mN EDTA) and 1 µl T4	
	polynucleotide kinase (10 U/µ1) were added and and	
	the tube was incubated at 37°C for 15 minutes.	
	Labelled olignucleotide was separated from	
15	unincorporated label by ethanol precipitation in the	
	presence of ammonium acetate, followed by ethanol	
	prodict to the time to the tim	
	pellet was dissolved in 50 µl of TE and 1 µl was	
	counted by emmission of Cerenkov radiation. The	
20	incorporation of ³² P by this method was 3-8 x 10 ⁶	
	counts per pmole of oligonucleotide. Ten µg of RNA	
	from the roots of both untreated and N-(amino-	
	carbonyl)-2-chlorobenzenesulfonamide-treated plants	
25	were annealed to 0.2 pmoles of oligonucleotide in a	
2.5	volume of 10 μl in 0.25 M KCl, 2 mM Tris-HCl pH 7.9	
	and 0.3 mM vanadyl ribonucleoside complex (BRL) at	
	37°C, 45°C and 55°C for 3 hours. To the annealed	
	RNA, 23.5 µl of primer extension mix (10 mM MgCl ₂ , 5	
	mM DTT, 20 mM Tris-HCl pH 8.3, 0.33 mM d(GATC) TP,	
30	100 μg/ml actinomycin D) and 0.5 μl (10 units) avian	
	reverse transcriptase (Life Sciences) were added and	
	the mixture was incubated for 45 minutes at 37°C.	•
	The nucleic acids in the reaction were precipitated	•
	with ethanol and dried. The pellet was dissolved in	•
35	3 µl of 0.1 M NaOH, 1mM EDTA and the solution was	•
	left at room temperature for 30 minutes to hydrolyze	
	-	

the RNA template. Six ul termination dye (Example 1) was added and the sample was heated at 80°C and 5 quick-cooled. The primer extension products were separated on a 6% denaturing polyacrylamide sequencing gel. A 110 bp long primer extension product was observed. predicting an untranslated leader of 68 bp. 10 To determine the 5' endpoint of the P6.1 gene. two fragments of the gene were subcloned for RNAse protection analysis. Both fragments span the first in frame ATG downstream by 40 bases (to a Nhe I site) and upstream by either 130 (a DraI site) and 300 (a 15 Spe I site) bases. Twenty µg of P622 DNA were digested with to completion with both Spe I and Nhe A separate aliquot of P622 was digested to completion with Dra I and Nhe I. The digestion products were separated by electrophoresis on a 5% 20 acrylamide gel, and 340 bp Spe I/Nhe I and 170 bp Dra I/Nhe I DNA fragments were cut out of the gel and recovered by electroelution. The DNAs were extracted with phenol:chloroform (1:1 v/v) and precipitated with ethanol. These fragments were subcloned into 25 the transcription vector Bluescript+ (BS+) (Stratagene). To accomplish this, 10 µg of BS+ was digested with Sma I and Xba I to subclone the Dra I/Nhe I fragment and Spe I and Xba I to subclone the Spe I/Nhe I fragment. BS(+) DNA was then extracted 30 with phenol:chloroform (1:1 v/v) and precipitated with ethanol. Ligations were performed at room temperature for 2 hours in volumes of 10 μ l. An aliquot of the ligation mixture was used to transform competent E. coli MV1193 using an X-gal selection. 35 Small scale plasmid preparations were performed on a number of white colonies and the DNAs were digested

with Eco R1 and Sac I. A colony containing a plasmid with the Dra I/Nhe I fragment in BS(+) was identified and designated P644. A colony containing a plasmid with the Spe I/Nhe I fragment in BS(+) was identified and designated P645. RNA probes complementary to the coding strands in both P644 and P645 were synthesized in the following reaction: 50 uCi 32P-UTP, 2 ul 5 X 10 transcription buffer (200 mM Tris-HC1 pH 7.5, 30 mM MgCl2, 10 mM spermidine), 0.5 μ l 0.2M DTT, and 0.5 μ l of either T3 polymerase (plasmid P645) or T7 polymerase (plasmid P644). Incubation was carried 15 out at 40°C for 1 hour. The DNA template was hydrolyzed for 15 minutes at 37°C by addition of 30 ul H-O, 1 ul RNAsin, 2.5 ul vanadyl ribonucleoside complex, 6 ul 5 X transcription buffer and 1 ul DNAse 1 (1 mg/ml) to the transcription reaction. The 20 reaction was extracted with an equal volume of phenol:chloroform (1:1 v/v). The RNA was precipitated once with ethanol in the presence of ammonium acetate and once with ethanol in the presence of sodium acetate. The pellets were dissolved in 25 ul of TE. Ten µg of RNA from 25 untreated and N-(aminocarbonyl)-2-chlorobenzenesulfonamide-treated plants were mixed with 1 x 106 cpm of each of the two probes in 30 ul of hybridization buffer (40 mM PIPES pH 6.7, 0.4 M NaCl, 1 mM EDTA). The mixture was then overlayed with 30 3.0

> RNAse T1 to 40 µg/ml and 2 µg/ml respectively in 10 35 mM Tris-HC1 pH 7.5, 5 mM EDTA, and 300 mM NaCl. Digestion was carried out at 30°C for 1 hour and

µ1 of mineral oil and hybridizations were carried out at 45°C for 16-24 hours. Single stranded RNA was selectively digested by adding 300 µ1 RNAse A and RNAses were inactivated by the addition of 20 μ 1 10%

- SDS and 50 pg of proteinage K followed by a 15 minute incubation at 37°C. The reaction mixture was extracted with phenol:chloroform (1:1 v/v) and the RNA hybrids were precipitated with 1 ml of ethanol after addition of 20 μg carrier of yeast tRNA. The pellets were dried and dissolved in formamide loading buffer. The samples were denatured at 90°C for 3 10 minutes and analyzed on a denaturing acrylamide gel. Protected fragments of 110 bp were observed in induced but not control RNA using both probes. These results agree with the predicted transcriptional 15 start site from the primer extension analysis. The sequence of the P6.1 gene 5' to its translation start site is shown in Figure 8. The arrow indicates
 - The 3' end of the gene was deduced from comparison of genomic and cDNA clone sequence data.

deduced transcription start site.

Construction of p614

20

- A 4.5 kb Hind III/Sal I genomic fragment from phage 2 containing the P6.1 petunia gene was
- 25 subcloned into pUCl18. 20 μg of the genomic phage 2 was digested with Hind III and Sal I, and the products separated by agarose gel electrophoresis. The 4.5 kb band containing the gene was isolated by electroelution as described earlier. Ten μg of
- pucils was digested to completion with Mind III and Sal I and the vector was then purified from the polylinker fragment by chromatography on Sepharose® CL-28 (Pharmacia). Vector and insert were ligated together in a volume of 10 µl overnight at 15°C, and
- 35 a portion of the ligation mixture was used transformed competent E. coli JM83. Aliquots of the

10

transformation mixture were plated out on LB plates containing 75 g/ml ampicillin which had been spread with X-gal and IPTG prior to plating of bacteria. Small scale plasmid prearations were performed on white colonies and DNAs were digested with Hind III and Sal I until a colony was found that contained the 4.5 kb Hind III/Sal I genomic fragment containing the petunia F6.1 gene. This plasmid was designated F614.

Construction of P654

Convenient restriction sites were introduced into the P6.1 petunia gene at the translation start and stop sites of the P6.1 coding region to use the regulatory regions from the inducible petunia gene to test if they could be made generally useful for expressing foreign coding regions in transformed plants. Site-directed mutagenesis was performed on P8.14 to introduce an New York.

- P614 to introduce an Nco I site was at the translation initiation ATG of the gene using the oligonuclectide 5-CGTTAGCCATGGTTATGCTTA-3'. The methods used to accomplish this mutagenesis were described in Example 1. The plasmid resulting from
- 25 the addition of an Nco I at the translation start site of the P6.1 gene fragment in P614 was designated P653. The plasmid P653 was further mutagenized using the oligonucleotide 5'-GCATATGCATAGATCTTATTGAATTCC-3' to introduce a Bgl II site at the translation stop
- 30 codon of the P6.1 gene. The resulting final plasmid construction, containing a petunia P6.1 gene with Nco I and Bgl II sites bounding the coding region of the P6 protein coding region, was designated P654 (Figure 9).

EXAMPLE 6

5 Isolation of the T2.1 Tobacco Gene

Isolation of cDNA T2

The procedures described for the isolation of the petunia cDNA clone P6.1 in Example 4 were

10 repeated using N. tabacum (Petite Havana SRI) as the starting plant material. Differential screening of the resulting tobacco cDNA library prepared using poly(A)* RNA from the roots of N-(aminocarbonyl)-2-chlorobenzenesulfonamide-treated tobacco plants identified a cDNA clone representing an N-(aminocarbonyl)-2-chlorobenzenesulfonamide-

inducible mRNA species. This clone was designated T2.

The insert from T2 was subcloned into the

vector pUCl19 as a single Eco RI fragment using
20 methods described in Example 4 for the sucloning of
the insert of the P6 cDNA clone. The resulting
plasmid containing the 1 kbp cDNA insert from cDNA
clone T2 in the Eco RI site of pUC 119 was called
T2.1 The same 1 kbp Eco RI cDNA fragment was also
25 cloned into the Eco RI site of the vector pBS (-).

with the resulting plasmid being designated T2.11.
A Northern blot of total RNA from the roots of untreated and N-(aminocarbony1)-2-chlorobenzenesulfonamide-treated tobacco plants was probed with

nick-translated T2.1 to determine the size of the corresponding T2 mRNA. The methods used for these procedures were described in Example 4. The T2.1 plasmid hybridized to an mRNA of 800 nt in RNA from the roots chemically treated plants, but not present

35 in control plants. This indicated that cDNA T2 represented an N-(aminocarbony1)-2-chloro-

15

benzenesulfonamide-inducible mRNA species, and the insert in the cDNA clone was full-length. 5 that the RNA appeared smaller than the cDNA clone suggested that T2 may contain some artifactual sequence generated during its cloning. The DNA sequence of T2 cDNA was determined by analyzing a set of deletions of T2 prepared as described previously. Examination of the sequence revealed that T2 contained a perfect inverted repeat from bases 11 to 164 and 518 to 671. Since the open reading frame begins past base 164, it was assumed that the first 164 bases were an artifact of cDNA synthesis and/or cloning that gave rise to a cDNA

larger than its corresponding mRNA. The predicted peptide encoded by the T2 cDNA contains the same number of amino acids as the petunia gene P6 and is 95% similar at the amino acid level. It was

20 therefore assumed that the T2 cDNA clone from tobacco represented a gene which is homologous to the petunia P6.1 gene.

Isolation of genomic clone T2.1

25 A genomic library was prepared from SRI tobacco as described in Example 4. A 32p-RNA probe was synthesized with T7 DNA polymerase using the cDNA insert of T2.11 as a template and and the resulting RNA transcript was used to screen the SRI genomic 30 library as described earlier. From this screening, a plaque was identified with homology to the T2 cDNA. This phage was plaque purified and designated phage #9. DNA purified from phage #9 was digested with the restriction enzymes Eco RI, Bam HI, and Sal I and the

35 resulting restriction fragments were separated by agarose gel electrophoresis and blotted to

nitrocellulose. The blot was then prehybridized and hybridized with nick-translated T2.11. Results of this blotting experiment revealed that the cDNA probe hybridized to a unique 5.0 kbp Bam HI/Eco RI fragment. This Bam HI/Eco RI fragment, believed to contain a complete copy of the T2 dene, was then cloned into the vector pUCl18 that had been digested 10 to completion with Bam HI and Eco RI. The resulting plasmid was called T217 (Figure 10). The gene contained within phage #9 was designated T2.1. The 5' end of the T2.1 mRNA was mapped by primer extension analysiz: The Oligonucleotide used 15 in this analysis was the same one chosen for analysis of the 5' end of the petunia P6.1 mRNA. This resulted in one mismatch relative to the tobacco gene positioned in the center of the oligonucleotide. Annealing of the primer was therefore performed at a 20 lower temperature for the tobacco mRNA (25°, 30°, and 35°C). Primer extension was then performed as described in Example 4. The primer extension product observed in this analysis was 110 bases long; exactly

the length of the extension product observed using 25 the petunia P6.1 mRNA as a template. This indicates that the 5' untranslated leader in the T2.1 mRNA was also 68 bp.

It is anticipated that those skilled in the art will be able to identify the promoter and downstream 30 regulatory regions of the T2 gene by following methods and procedures described in Example 4. Later examples teach the use of such regulatory regions.

EXAMPLE 7

Construction of Recombinant Genes Whose Expression

are Controlled by 2-1 Corn Promoter and 3' Downstream
Region

Construction of plasmids pJE 514 and pJE 516

Plasmids p484-1(Nco I) and p484-62 (Bgl II)

10 from Example 1, which contained convenient
restriction sites at the start and stop sites
respectively of the 2-1 structural gene were used to
create a new 2-1 gene from which the native coding
sequence could be easily removed and replaced with
15 foreign structural gene. Introduction of such a

recombinant gene into transgenic plants should place expression of the foreign coding region under the control of substituted benzenesulfonamides.

To construct this new 2-1 gene, pJE 484-1 (Nco I) (Figure 5) was digested to completion with Eco RI and Sma I, and 10 µg of digested DNA was subjected to electrophoresis on a 1% agarose gel overnight at 20v. The gel was stained with ethidium bromide and the DNA was visualized on a long wave UV trans-

25 illuminator. A small trough was cut in the gel just ahead of the desired 7.5 kb insert fragment. The DNA was electroeluted into this trough at 300 V and buffer containing the DNA was transferred to a microcentrifuge tube. The purified DNA fragment was

30 then extracted with an equal volume of phenol:chloroform (1:1 v/v), ethanol precipitated and resuspended in 10 μl of H₂O. The plasmid pJE 484-62(Bgl II) was digested with Eco RI and Nco I, and a 1.3 kb fragment was gel-purified in the manner

35 described above. The 7.8 and 1.3 kbp DNA fragments were ligated together in 10 μl of 1% ligase buffer as

10

15

20

described in earlier examples and the ligation
products were used to transform competent E. coli
JM83 Cells Small scale plant's
JM83 cells. Small scale plasmid preparations were
performed on transformed colonies and diagnostic
restriction enzyme digestions were performed on
individual colonies until one was found that
contained a copy of the 2-1 corn gene with the added
Nco I and Bgl II sites at its respective translation
start and stop sites. This construction was
designated pJE 514.
The coding sequence chosen to replace the 2-1
coding sequence of pJE 514 was β-glucuronidase
(referred to as GUS) [Jefferson R., Proc. Natl. Acad.
Sci. USA. 83: 8447-8451, 1986). A GUS coding
Sequence was included (
sequence was isolated from the plasmid pJJ 3892 as a
1.8 kbp Nco I/Xba I fragment. The identical 1.8 kbp
Nco I/Xba I fragment is available in the plasmid
pJJ 3431 (ATCC accession number 67884, and described
in Example 9), and thus pJJ 3431 can be substituted
for pJJ 3892 in this Example. To this end pJJ 3892
Was discarded to I this end pJJ 3892
was digested to completion with Xba I and the
resulting 5' overhangs were blunted using the Klenow
fragment of DNA polymerase I as described in earlier
- Curiter

25 examples. After extraction with phenol:chloroform (1:1 v/v) and ethanol precipitation, the DNA was digested to completion with Nco I and the resulting DNA fragments were separated by agarose gel

30 electrophoresis. A 1.9 kbp DNA fragment corresponding to the GUS coding region was recovered from the gel and ligated with pJE 514 that had been digested to completion with Bgl II, blunt-ended with Klenow fragment of DNA polymerase I and then digested to completion with Nco I. An aliquot of this

35 ligation mixture was used to transform E. coli HB101

30

and individual transformants were analyzed until one was found that contained the GUS coding sequence in place of the 2-1 structural gene. This plasmid was designated pJE 516 (Figure 11).

Construction of plasmid pDuPE2

Plasmid pJE 516 was used as the starting
material to produce a deletion series consisting of a
GUS gene/2-1 3' downstream region fusion whose
expression is regulated by progressively smaller 2-1
promoter fragments. The deletion series was
generated by linearizing 40 µg of pJE 516 DNA with 25
units of Hpa I restriction endonuclease in 20 mM KCI,
10 mM Tris-HCI pH 7.4, 10 mM MgCl₂ and 1 mM DTT in a
final volume of 100 µl. The reaction was incubated at
17°C for 3 hours, and DNA was extracted with am gual

volume of phenol:chloroform (1:1 v/v) and
20 precipitated with ethanol. The linearized DNA was
recovered by centrifugation and dried in vacuo.

The DNA pellet was resuspended in 180 μ l of $\rm H_{2}O$ and 30 μ l of 10X Bal 31 buffer was added (final concentrations in the reaction were 20 mM Tris-HCl pH 8, 12 mM MgCl $_{2}$, 12 mM CaCl $_{2}$ and 300 mM NaCl $_{3}$. The Bal 31 digestion was carried out as recommended by the manufacturer (Bethesda Research Labs) using 2 units

of Bal 31. This mixture was incubated at 30°C for various time intervals (e.g. 0, 2.5 or 5 minutes), and the reaction in each aliquot was stopped by

adding 50 µl of 100 mM EDTA, pH 7.6. The DNA was then extracted twice with 100 µl of phenol, twice with 100 µl of CHCl₃, then precipitated with 2.5 volumes of ethanol. Bal 31 digested DNA was recovered by

35 centrifugation and dried in vacuo.

The dry DNA pellet was dissolved in 100 ul of Sal I buffer (150 mM NaCl 10 mM Tris-HCl pH 8, 10 mM ${\rm MgCl}_{2}$ and 10 mM β -mercaptoethanol) and digested with 50 units of Sal I for 4 hour at 37°C. The reaction was extracted with phenol:chloroform (1:1 v/v) and ethanol precipitated as above. The ends of the DNA were rendered blunt using the Klenow fragment of DNA Polymerase I as follows: DNA was dissolved in 60 ul 10 of 66 mM Tris-HCl pH 7.6, 6.6 mM MgCl2, 52 mM NaCl, 1 mM β-mercaptoethanol, 0.5 mM dNTPs and 10 units of Klenow. The reaction was incubated at room temperature for 2 hours. The DNA was then 15 fractionated by electrophoresis in a 0.7% low melting agarose gel. The gel was stained with 1 µg/ml ethidium bromide solution, and a gel piece containing the DNA fragment of the desired deletion length was excised from the gel under UV illumination. The gel piece was frozen at -80°C for 20 minutes, thawed, 20 crushed with a pipette tip, and centrifuged for 30 minutes in a microcentrifuge. The aqueous solution was transferred to a fresh tube, adjusted to a final concentration of 0.3 M sodium acetate and 2.5 volumes of ethanol were added. The precipitated DNA 25

- was recovered by centrifugation, dissolved in 20 ul of water and was self-ligated (recyclization). Ligation reactions were performed in 50 mM Tris-Cl pH 7.8, 10 mM MgCl2, 20 mM DTT and 1 mM ATP. The ligation reaction was carried out at room temperature 30
- for 8 hours, and diluted five fold with water prior to using it to transform of competent E. coli HB101 cells. Aliquots of the transformation mixture were spread on LB plates containing 50 µg/ml of amp and 35
- plates were incubated overnight at 37°C.

Individual amp resistant colonies were picked and grown up at 37°C with vigorous shaking in 2 ml of

- 2XTY containing 50 ug/ml amp. Small scale plasmid preparations were performed on the bacteria and aliquots of the DNAs were digested to completion with Nco I and Xho I. The resulting DNA fragments were analyzed by 1.5% agarose gelelectrophoresis to
- 10 determine the size of the 2-1 promoter fragment remaining in each plasmid. Results from the analysis showed that one clone, designated pDuPE2 contained the GUS construction of pJE 516, operably linked to a 900 bp 2-1 promoter fragment (relative to the
- translation start site of the 2-1 gene). 15

Construction of plasmids pDuPI8 and pDuPI9

- The Bal 31 digestion protocol used to create pDuPE2 was repeated using the plasmid pDuPE2 as the starting material to create progressively shorter 2-1 20 promoter fragments. DNA was first linearized with Xho I, followed by Bal 31 digestion at different time intervals (from 2-5 min). The Bal 31-digested DNA was extracted with phenol:chloroform (1:1 v/v).
- ethanol precipitated, and 5' ends of the DNA were 25 filled-in using Klenow fragment. DNA was then further digested with Bam HI to excise the entire remaining 2-1/GUS construction from the pBS(+) vector. The Bam HI digested DNA fragments were
- separated by electrophoresis in a 1% low-melting agarose gel and the DNA fragments containing the deleted constructs were extracted as described above and ligated into the Bam HI-Sma I sites of the vector pBluescript (S/K)+ vector (Stratagene). The ligation
- mixture was diluted four fold with with H2O, and 35

	aliquots of the transformation reaction were spread
5	onto LB plates containing 50 µg/ml amp and incubated
-	overnight at 37°C. Small scale plasmid preparations
	were performed on amp-resistant colonies and DNAs
	were digested to completion with Nco I and Xho I. A
10	series of clones containing 2-1 promoter fragments
	ranging in size from 500 to <100 bp was chosen from
	these colonies. The designated names of thee
	constructions and the length of the 2-1 promoter
	fragment in each is shown in Table 1.

TABLE 1

15

20

Construction	
Designation	Promoter Length (bp)
pDuPE2	~900
pDuPI8	421
pDuPI9	226

EXAMPLE 8

25 Construction of Recombinant Genes Whose Expression is Regulated by the 2-2 Corn Promoter and Various 3' Downstream Regions

Construction of plasmid pHPH201(+)

30 Plasmid pRAJ275 (available from Clontech Laboratories, Inc. 4055 Fabian Way, Palo Alto, CA 94303) served as a source for an E. coli β-glucuronidase (GUS) gene in this construction. The GUS coding region in pRAJ275 has a unique Nco I site 55 positioned at the initiator ATC codon of the protein coding sequence.

Genomic subclone 2-2#4-17 (Example 2) (320 mg) was partially digested with Nco I for 1 hour at 37°C using 0.5 units of enzyme per microgram of plasmid DNA. The digestion was stopped by addition of Na,EDTA to a final concentration of 20 mM and DNA was ethanol precipitated in the presence of 0.3 M sodium acetate, pH 6.0. The partially digested plasmid was dissolved in 260 μl of TE, pH 8.0 and 40 μl of 10 electrophoresis tracking dye. The DNA was loaded into 4 cm X 1 cm X 2 mm wells of a 2 mm thick 5 % polyacrylamide gel in 1 X TBE buffer and sujected to electrophoresis at 325 volts for 4 hours. A 1.68 kbp Nco I fragment was recovered from each lane of the 15 gel as described earlier. One half of the purified Nco I fragment was ligated overnight in a total volume of 10 µl with 0.5 µg of pRAJ275 that had been cut to completion with Nco I and dephosphorylated. The ligation mixture was diluted to 50 μ l with H_2O . 20 and 3 µl of the dilution was used to transform 60 µl of competent E. coli HB101 cells. Aliquots of the transformation reaction were spread onto LB plates containing 50 µg/ml amp and plates were incubated 25 overnight. Small scale plasmid preparations were performed on amp-resistant colonies until one was found that contained 1.68 kbp Nco I promoter fragment ligated into pRAJ275 such that it was operably linked to the 5' end of GUS gene. This plasmid was called pHPH201(+). 30

Construction of p2-2 Hind III 3' end

A construction containing the 3' end of the 2-2
gene that is generally useful in preparing

recombinant genes whose expression is controlled by

35 recombinant genes whose expression is controlled by substituted benzenesulfonamides was prepared. 7.0

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Genomic subclone 2-2#4-11 (Figure 4A) was digested to completion with Hind III. The 5' overhang was filled-in using the Klenow fragment of DNA polymerase I, and the DNA was extracted sequentailly with a equal volumes of phenol:chloroform (1:1 v/v) and chloroform. The DNA was ethanol precipitated. collected by centrifugation and redissolved in TE pH 8.0. The vector pUC18 was cut to completion with Sac I and Kon I and the resulting 3' overhangs were removed using the Klenow fragment of DNA polymerase I. The DNA was extracted with phenol:chloroform (1:1 v/v), precipitated with ethanol and redissolved in TE pH 8.0 as described above. The blunt-ended Hind III digestion products of genomic 2-2#11 (0.6 μ g) were then ligated with 0.45 µg of the blunt-ended pUC 18 DNA overnight at 16°C. The ligation mixture was diluted to 50 µl with H2O, and 1 µl of the dilution was used to transform 20 ul of competent E. coli HB101 cells. Aliquots of the transformation reaction were spread onto LB plates containing 50 µg/ml amp and plates were incubated overnight. Small scale plasmid preparations were performed on amp-resistant colonies and the resulting DNAs were digested with Eco RI and Bam HI until a colony was found that contained the 2.3 kbp Hind III fragment of genomic subclone 2-2#11 blunt-ended into the Kpn I/Sac I sites of pUC18. This plasmid construction was

called p2-2 Hind III 3' end. Construction of plasmid pHPH102

Plasmids p2-2 Hind III 3' end and the vector pMspFr (ATCC accession number 67723) were both digested to completion with Eco RI and Hind III. Following dephosphorylation of pMspFr, 1.6 µg of

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		vector was ligated overnight with 0.38 µg of Eco	
	5	RI-Hind III digested p2-2 Hind III 3' in a final volume of 10 μ l. The ligation was diluted to 50 μ l with H_2O and 1 μ l of the dilution was used to	•
	10	transform 60 µl of competent HB101 cells. Alignots of the transformation mixture were spread onto LB plates containing 100 µg/ml of both spectinomycin and streptomycin (spec/strep) and plates were incubated overnight at 37°C. Small scale plasmid preparations were performed on spec/strep-resistant colonies and the resulting DNAs were digested with Eco RI and Hind III until one was found that contained the desired downstream sequences of the 2-2 gene on a 2.3 kbp Eco RI-Hind III fragment. The resulting plasmid was	:
4		Construction of plasmid pHPH 220	
	20	The plasmid pHPH 102 was cut to completion with Xho I and the resulting 5' overhang was filled-in with Klenow fragment of DNA polymerase I. The blunt-ended DNA fragment was dephosphorylated as described in Example 1 and then cut to completion	
	25	with Hind III. Plasmid pHPH201(+) was partially	

cleaved with Eco RI by digesting it with Eco RI at 37°C for 90 minutes using 0.85 units of enzyme per microgram of DNA. Eco RI was inactivated by heating the digestion mixture to 70°C for 10 minutes, and the resulting 5' overhang was filled-in with Klenow 30

fragment as described above. This DNA was then digested to completion with Hind III and 2.1 µg of the resulting DNA was ligated overnight in a final volume of 15 µl with 0.8 µg of Hind III cut pHPH102 35

that had been blunt-ended at its unique Xho I site. The ligation mixture was diluted to 60 µl with HaO

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and 1 µ1 was used to transform 80 µ1 of competent E. <u>Seli HB101 cells</u>. Aliquots of the transformation mixture were spread onto LB plates containing 100 µg/ml of both spec/strep and plates were incubated overnight at 37°C. Small scale plasmid preparations were performed on spec/strep-resistant colonies until one was found that contained the 3.6 kbp Hind III-Eco RI fragment from pHPH201(+) (consisting of the 1.7 kbp 2-2 promoter/GUS coding region fusion) operably linked to the 2.3 kbp or downstrean sequence originating from the 2-2 gene in the vector pMSP^TK. This plasmid was called pHPH 220 (Figure 12).

Construction of plasmid pIn 2-2(3.9)

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Two and a half µg of DNA from genomic clone 2-2 #4 (Example 2) was digested to completion with Sal

I. One µg of pUC18 DNA was also digested to

20 completion with Sal I. The DNAs were extracted with equal volumes of phenol, phenol:chloroform (1:1 y/y) and chloroform. The DNA was then precipitated with ethanol in the presence of sodium acetate. A ligation reaction was carried-out overnight at 16 C 25 with a 3:1 ratio of genomic 2-2#4 DNA to pUC18 in a volume of 10 µl. The ligation mixture was diluted 5 fold with water and an aliquot of the ligation mixture was used to transform competent E. coli DH5 α cells. Aliquots of the transformation reaction were plated on LB agar plates containing 50 µg/ml amp, 25 30 mM IPTG and 40 µg/ml X-Gal. Plasmid DNA from individual white colonies was prepared and digested to completion with Sal I. A clone was identified

which contained the 3.9 kbp Sal I fragment from the 35 2-2#4 DNA which encompassed a region of the 2-2 gene extending from 3.6 kbp 5' from the translation start

of the 2-2 protein to 180bp inside the coding region of the 2-2 protein. This plasmid was designated pIn 2-2(3.9).

Construction of pTDS130

Twenty-five µg of pJE516 was digested to completion with Nco I and Xho I. The DNA fragments were dephosphorylated with 24 units of calf intestinal alkaline phoshatase for 40 minutes at 37°C. Fifty µg of plasmid pIn 2-2(3.9) DNA was cut to completion with Pvu I and dephosphorylated as described above, precipitated with ethanol in the 15 presence of 0.3 M sodium acetate and resuspended in TE pH 8.0. This DNA was then digested to completion with Xho I. Partial Nco I cleavage of the resulting pIn 2-2(3.9) DNA was performed by digesting the Xho I

- digested DNA sample with 1 unit of Nco I at 37°C and removing 1/4 of the digestion mixture at 15 minute intervals. The Nco I digestion was stopped in each time point by addition of EDTA to a final concentration of 40 mM. The DNAs were extracted sequentially with equal volumes of phenol.
- 25 phenol:chloroform (1:1 v/v) and chloroform. DNA was precipitated with two volumes of ethanol, recovered by centrifugation and redissolved in 10 µl of TE, pH 8.0. Small aliquots of DNA from each digestion time were analyzed by agarose gel electrophoresis to find the digestion that contained the bitchest arms.
- 30 the digestion that contained the highest amount of the desired 1.9 kbp Xho I-Nco I promoter fragment. A total of 0.5 µg of partially digested DNA was ligated with 0.18 µg of pJE 516 DNA overnight at 16°C. The ligation reaction was heated at 70°C for 10 minutes,
- 35 diluted 5 fold with water, and 2 µl of the dilution was used to transform 100 µl of competent E. coli

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HB101. Aliquots of the transformation mixture were plated on LB syst plates containing 50 µg/ml amp and allowed to grow overnight at 37°C. Plasmid DNA prepared from amp-resistant colonies were analyzed by restriction endonuclease digestions until one was identified that contained the 1900 bp Xho I/Nco I Promoter fragment of the 2-2 gene operably linked to the GUS/2-1 3' end downstream region fusion in the plasmid pJE516. This clone was designated pTDS130 (Figure 13).

Construction of plasmid pTDS133

Plasmid pTDS130 was cut to completion with both Eco RI and Xho I and the enzymes were inactivated by heating the reaction at 40°C for 20 minutes in the presence of 0.02% diethylpyrocarbonate (DEP). Excess DEP was destroyed by heating at 70°C

20 for 10 minutes, and 5' overhangs in the DNA were filled-in with the Klenow fragment of DNA polymerase I.

The DNA was extracted sequentially with equal volumes of phenol, phenol:chloroform (1:1 v/v) and chloroform followed by ethanol precipitation in the presence of sodium acetate. The DNA was then recircularized by subjecting it to overnight self-ligation. The ligation reaction was diluted five fold with water and 2 µl of the mixture was used to transform 100 µl of competent E. coli HBlO1. Aliquots of the transformation mixture were plated on LB agar plates containing 50 µg/ml Amp and allowed to grow overnight at 37°C. Small scale plasmid

preparations were made from individual amp resistant 35 colonies and analyzed by restriction endonuclease digestions until one was identified that contained

the 465 bp Eco RI/Nco I promoter fragment of the 2-2 gene was operably linked to the GUS/2-1 fusion in the

5 plasmid pTDS130. This plasmid was designated pTDS133 (Figure 13).

Construction of plasmid pTDS134 and pTDS136

Ten µg of pTDS133 DNA and 10 µg of the vector pBluescript SK(+) DNA were digested to completion with Bam HI. Vector DNA was dephosphorylated as described in earlier examples. Both DNAs were extracted with phenol:chloroform (1:1 v/v) and precipitated with ethanol. The digested pTDS133 and 15 pBluescript SK(+) were ligated together at a 3:1

molar ratio (insert:vector) in a final volume of 10 ul overnight at 16°C. The ligation mix was diluted five fold with water and 2 µl of this dilution was

used to transform 100 µl of competent E. coli HB101. 20 Small scale plasmid preparations were made from individual amp resistant colonies and analyzed by restriction endonuclease digestions until one was identified that contained the 3.4 kbp Bam HI fragment from pTDS133 cloned into the Bam HI site of

- 25 pBluescript S/K(+) in an orientation such that the 2-2 promoter was immediately adjascent to the Sma I site of the vector's polylinker. This plasmid construction was designated pTDS134 (Figure 14). A second colony containing the same 3.4 kbp Bam HI
- fragment cloned in the opposite orientaton such that 30 the 2-2 promoter was immediately adjascent to the Spe I site of the vector's polylinker was also identified. This plasmid construction was designated pTDS136.

Construction of plasmid pTDS231

The plasmid pDH51 was disclosed by Maciej

5 Pietrzak et al. and is described in Nucleic Acids Research, 14: 5857-5868 (1986).

Ten µg of pHPH201(+) DNA was digested to completion with with both Eco RI and Pvu I for two hours at 37°C, and the resulting 5' overhangs were

- 10 filled-in with Klenow fragment of DNA polymerase I.

 Ten µg of pDH51 DNA was digested to completion with
 Pst I and Nco I, and the resulting 5° and 3°
 overhangs were blunted with Klenow fragment of DNA
 polymerase I. The DNA samples were extracted
- 15 sequentially with equal volumes of phenol, phenolichloroform (1:1 v/v) and chloroform followed by ethanol precipitation. The blunt-ended pDH51 was then digested to completion with Bam H1 and dephosphorylated. The pDH51 DNA (0.25 µg) was
- 20 ligated overnight at 16°C with 0.75 μg of digested pHPH201(+) DNA in a final volume of 10 μl. The ligation reaction was heated for 10 minutes at 70°C and then diluted five fold with water. A 2 μl aliquot of the diluted ligation mixture was used to transform 100 μl of competent HB101 cells. Aliquots
- of the transformation mixture were plated on LB agar plates containing 50 µg/ml amp and allowed to grow overnight at 37°C. Small scale plasmid preparations were made from individual amp resistant colonies and 30 analyzed by restriction endonuclease digestions until
- one was identified that contained a plasmid consisting of the 465 bp Eco RI/Nco I 2-2 promoter/GUS fusion from pHPH201(+) operably linked to the 3' end fragment derived from the CaMV 35s
- 35 transcript in the plasmid pDH51. This clone was designated pTDS231 (Figure 15).

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Construction of 2-2 promoter deletions of pTDS130

Plasmid pTDS130 contains a unique Eco RI site that cleaves the 2-2 promoter 465 bp 5' to the ininiatior ATG codon of the 2-2 protein. This Eco RI site was cleaved to linearize pTDS 130 and provide a convenient starting point for the generation a of Bal

- 31 deletions of the promoter in this DNA construc-10 tion. The procedure used to create the 2-2 promoter deletion series from this Eco RI site was described in Example 7. All deletions were subcloned into pBluescript (SK)+. A series of cDNA clones with shorter 2-2 promoter fragments regulatng GUS
 - expression (increasing Bal 31 digestion) was selected from the deletions series generated above. The plasmid constructions selected for analysis are shown in Table 2 with the length of the 2-2 promoter
- 20 fragment remaining from the 5' end of the promoter to the translation start site in each construction. Promoter fragment lengths were determined by DNA sequence analysis of each construction.

25 TABLE 2

	Construction Name	Promoter Length (bp)
	pTDS133	465
30	pTDS134	450
	pDuPM17	248
	pDuPN27	208
	pDuPN4	150
	pDuPN7	120

35 The DNA sequence of the 2-2 pr moter region with the locations of the start sites of each of the

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promoter fragments driving the expression of GUS in the various constructions is given in Figure 14.

Construction of pDuPS22

A construction consisting of a recombinant gene encoding a sulfonylurea-resistant form of acetolactate synthase (ALS) under the transcriptional control of an inducible promoter fragment from the corn 2-2 gene was prepared. The details of the particular embodiment of such a construction presented here represents but one of any number of methods by which such a recombinant gene might be accomplished. It is expected that those skilled in the art will be able to make such recombinant gene using the sulfonylurea-resistant ALS gene contained in pAGS148 (ATCC accession number 67124) and any

number of 2-2 promoter fragments whose use is taught 20 in this work.

The construction pUC119/HRA was made using the plasmid pAGS148 as the starting material (ATCC accession number 67124, and described in detail in European patent application 0257993). pAGS148 was 25 digested to completion with Eco RI and the 1.38 kbp Eco RI fragment containing the translation start site of the ALS protein, was subcloned into the Eco RI site of the vector pUC119. This construction was designated pUC119/AGS. The plasmid pUC119/AGS was 30 digested to completion with Bbv I and the 5 overhangs of the resulting fragments were blunted with the Klenow fragment of DNA polymerase I. These blunted fragments were separated by agarose gel electrophoresis and the 1.2 kbp fragment was purified 35 from the gel. Bam HI linkers (New England Biolabs, catalog #1017) were added to the fragment which was

	then subcloned into the Bam HI site of pUC119 to
	yield the plasmid pUC119/Bbv I.
5	The plasmids pUC119/Bbv I and pAGS148 were
	digested to completion with Bst EII and Pst I and the
	resulting fragments were separated by gel
	electrophoresis. The 4.58 kb BstE II/Pst I fragment
	from pUC119/Bbv I and the 2.45 kb Bst EII/PstI
10	fragment from pAGS148 were purified from the gels and
	ligated together to yield the plasmid pUCl19/HRA.
	Mutations were made in the tobacco SurA gene to
	change amino acid number 194 from proline to alanine
	and amino acid number 571 from trypotphan to leucine
15	as described by Bedbrook et al. in European patent
	application 0257993. The 1.42 kbp Nco I/Bgl II
	fragment corresponding to nucleotides 533-1952 of the
	SurA gene was excised by restriction endonuclease
	digestion and used to replace the corresponding
20	region in the pUC119/HRA to yield the plasmid pUCAD.
	The plasmid pTDS130 was digested to completion
	with Nco I. The 5' overhangs of the Nco I sites were
	partially filled-in with the Klenow fragment of DNA
	polymerase I by using only dCTP and dGTP as
25	nucleotides in the Klenow reaction. The remaining
	nucleotides of the overhangs that were not filled-in
	were removed by digestion with mung bean nuclease and
	the resulting blunted DNA fragments were separated by
	gel electrophoresis. A unique 450 bp DNA fragment
30	was isolated from the gel and ligated together with
	equimolar amounts of pUCAD that had been digested to
	completion with Bam HI and rendered blunt-ended by
	digestion with mung beam nuclease. The resulting
	plasmid, containing an ALS gene encoding a
35	sulfonylurea herbicide-resistant form of the enzyme

under the transcriptional control of a 450 bp

inducible 2-2 promoter fragment was designated

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EXAMPLE 9

Construction of Recombinant Genes Whose Expression is Regulated by the 5-2 Corn Promoter

10 Construction of pMC 710

The 2-1 promoter fragment in the construction DJE 516 was removed and replaced with a 5-2 promoter. To this end, pJE 516 was digested to completion with Sst II and the resulting 3' overhang was removed using T4 DNA polymerase. This DNA was then digested to completion with Nco I and the DNA fragments were separated by agarose gel

electrophoresis. The 3.8 kbp band corresponding to the GUS/2-1 3' end fusion from pJE516 was cut out of

- the gel and recovered as described earlier. The plasmid pMC 75.5 was digested to completion with Xho I and the resulting 5' overhang was filled-in using the Klenow fragment of DNA polymerase I. This DNA was then digested to completion with Nco I and dephosphorylated. The resulting DNA was ligated to
- the 3.0 kbp Nco I-blunt DNA fragment from pJE516. An aliquot of this ligation mixture was used to transform competent <u>E. Coli</u> HB101 and individual transformants were analyzed until one was found that
- 30 contained the 5-2 promoter operably linked to the GUS/2-1 3' end fusion in the vector pBS(-). This construction was designated pMC715.83 (Figure 17).

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EXAMPLE 10

Construction of a Chimeric Gene Whose Expression is

5 Controlled by The 218 Corn Promoter

The plasmid pMC791 (Example 4) was subjected to partial digestion with Afl III. The partially digested pMC791 was then digested to completion with Sma I. The digestion products were separated by gel electrophoresis and a 1.4 kbp Afl III/Sma I DNA fragment was isolated.

The plasmid pJE516 was digested to completion with Sal I and the resulting 5' overhang was filled in using T_A DNA polymerase. The DNA was then

- digested to completion with Nco I, dephosporylated 15 and ligated with an equimolar amount of the gel-purified 1.4 kbp Afl III/Sma I fragment from pMC791. An aliquot of the ligation mixture was used
- to transform competent E. coli HB101 cells. Aliquots 20 of the transformation mixture were spread on LB agar plates containing ampicillin and the plates were incubated overnight at 37°C. Plasmid DNA prepared
- restriction endonuclease digestions until one was 25 identified that contained the 1.4 kbp Sma I/Afl III promoter fragment of the 218 gene operably linked to the GUS/2-1-3' end fusion in pJE516. This plasmid

from amp-resistant colonies was analyzed by

was designated pMC7113 (Figure 18).

EXAMPLE 11

Construction of Recombinant Genes Whose Expression are Regulated by Petunia P6.1 Gene Promoter Fragments and Various 3' Downstream Regions

35 Construction of P655, P657, P658, and P660

Construction of P655

The reporter gene used for fusions was β-glucuronidase from E. coli as discussed in earlier

		examples. The source of this gene was the plasmid
		pJJ3431 (ATCC accession number 67884), which contains
	5	a GUS coding region fused to the 35S CaMV promoter
í		region and the octopine synthase 3' end in pUC118.
		The regulatory regions from P6 gene were substituted
;		into pJJ3431 in a stepwise fashion: first the 35S
		promoter was replaced with the no
	10	promoter was replaced with the P6 gene 1 promoter,
		then the octopine synthase (OCS) 3' end was replaced with the P6 gene 1 3' end.
		The 35S promoter region was removed from
		pJJ3431 by digesting 10 µg of the plasmid with Eco RI
		and filling-in the resulting 5' overhang with Klenow
	15	fragment. After extraction with phenol:chloroform
		(1:1 v/v) and precipitation with ethanol, the DNA was
		restricted with Nco I and the products were separated
		by agarose gel electrophoresis. A 5.8 kbp DNA
		fragment corresponding to the GUS/OCS 3'end fusion in
	20	pUCl18 was isolated by placing the gel slice
		containing this fragment in a dialysis bag with 500
		μl of 1X TAE buffer and electroeluting the DNA from
		the agarose. The eluted DNA was extracted with
		phenol:chloroform (1:1 v/v) and precipitated with
	25	ethanol. The mutagenized petunia P6 gene 1 promoter
		region containing a unique Nco I site was purified by
		digesting 10 µg of the plasmid construction P653
		(Example 4) to completion with Nco I and Sma I and
	30	gel purifying the 1.3 kbp P6 promoter fragment as
	•	previously described. Equimolar amounts of this 1.3
		kbp promoter fragment and the GUS/OCS 3' end fragment
:		were ligated overnight at 15°C in a volume of 10 μ l.
		The ligated DNA was used to transform competent E.
		coli JM83 and aliquots of the transformation mixture

35 were plated on LB containing 75 μg/ml amp. Small scale plasmid DNA preparations from amp-resistant

	colonies were evaluated by digestion with Nco I and
	Bam HI until a colonoy containing a plasmid with the
5	1.3 kbp P6 gene 1 mutagenized promoter fragment
	operably linked to the GUS/OCS fusion of pJJ 3431 was
	found. This plasmid was designated P655 (Figure 19).
	Construction of P657
10	In the construction P655, the petunia P6/GUS
	fusion was operably linked to an OCS 3' end at an
	Xba I site. In order to replace the OCS 3' end
	fragment in P655 with the mutagenized P6 gene 3' end
	in P654, it was necessary to first partially digest
15	P655 with Xba I as there was an Xba I site in the
	polylinker region of P655 in addition to the site of
	the OCS 3' end fusion. Due to a relatively inactive
	lot of Xba I, it was possible to generate partially
	cut molecules by digesting 10 µg of P655 DNA with 30
20	units of enzyme for 1 hour. After checking for
	partial digestion by agarose gel electrophoresis, the
	5' overhang of the Xba I site was filled in with
	Klenow fragment of DNA polymerase I. The DNA was
	extracted with phenol:chloroform (1:1 v/v), ethanol
25	precipitated, redissolved and digested to completion
	with Hind III. The products of this digestion were
	separated by agarose gel electrophoresis and the
	desired DNA fragment corresponding to P655 without
	the OCS 3' end was purified from the gel.
30	The 3' end of the P6.1 gene was isolated by
	digesting the plasmid P654 to completion with Bgl II
	and filling-in the resulting 5' overhang with Klenow
	fragment. The DNA was extracted with

phenol:chloroform (1:1 v/v), precipitated with

ethanol, redissolved, and digested to completion with Hind III. The resulting products were separated by

agarose gel electrophoresis, and the 2.2 kbp fragment containing the P6.1 gene 3' end was excised from the

- gel and purified as described earlier.
 - The 2.2 kbp P6 3' end fragment was ligated with the purified Xba I fragment of P655 from above overnight at $15^{\circ}C$ in a final volume of $10~\mu l$. An aliquot of the ligation reaction was used to
- transform competent E. <u>coli</u> JM83 cells. Small scale plasmid preparations from individual ampicillin resistant colonies were analyzed by digestion with Hind III and Bam HI until one was found that contained the P6.1 3' end operably linked to the P6.1
- promoter/GUS fusion. This plasmid was designated P657 (Figure 19).

Construction of P658

- In order to map potential regulatory regions in the promoter of P6.1, a 1 kb deletion was made in the promoter fragment of the P657 construction, leaving a
 - 300 bp P6.1 promoter fragment operably linked to a GUS/P6.1 3'downstream fragment. Ten µg of P657 was digested to completion with Xba I and Spe I. The
- 25 resulting 5' overhangs were filled-in with Klenow fragment and the products were separated by agarose gel electrophoresis. The 7.6 kb fragment (P657 with 1 kb of the 5' end of the promoter deleted) was
- recovered from the gel by electroelution, extracted
 30 with phenol:chloroform (1:1 v/v) and precipitated
 with ethanol. The DNA was ligated back to itself
- overnight at 15°C in a 10 µl ligation reaction. An aliquot of the ligation mixture was used to transform competent E. coli JM83. Plasmid DNA from individual
 - amp resistant colonies was digested with Hind III and Bam HI until a col ny containing the desired plasmid was found. This c lony, containing a GUS/OCS 3' end fusion operably linked to a 300 bp P6.1 pr moter fragment was designated P658 (Figure 19).

Construction of P660

A construction consisting of a GUS/OCS 3.

downstream region fusion operably linked to a 600 bp
P6.1 promoter fragment was prepared. A convenient
Eco RI site 600 bp upstream of the initiating codon
ATG was used to generate the 600 bp promoter

fragment. However, since 2 Eco RI sites are found in the 3' downstream region of the P6.1 gene, a promoter deletion was made in the plasmid P655 and the OCS 3' end was replaced with the 3' downstream region from the P6.1 gene.

Ten µg of P655 was partially digested with Xba I, extracted with phenol:chloroform (1:1 v/v) and precipitated with ethanol. The DNA was then digested to completion with Eco RI and the products separated.

by agarose gel electrophoresis. The 6.4 kbp DNA fragment corresponding to P655 lacking 700 bp from

the 5' end of the P6.1 promoter was purified, and the 5' overhangs were filled-in with Klenow fragment. The DNA was extracted with phenol:chloroform (1:1 v/v) and precipitated with ethanol. The 6.4 kbp

25 fragment was ligated to itself overnight at 15°C in a volume of 10 μl. An aliquot of the ligation mixture was used to transform competent E. coli JM83 cells. Plasmid DNA from individual amp resistant colonies was digested with Hind III and Bam HI until a colony

30 containing the desired 3.2 kb Hind III/Bam HI fragment was found, diagnostic of the presence of a 600 bp promoter fragment in the construction. This plasmid was designated P659 (Figure 20).

To replace the OCS 3' end of P659 with the 3' 35 end of the P6.1 gene, 10 µg of P659 DNA was first partially digested with Xba I. The 5' overhang was

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filled-in with Klenow fragment, and the blunt-ended
DNA was extracted with phenol:chloroform (1:1 v/v)
and precipitated with ethanol. The DNA was then
digested to completion with Hind III and the
resulting DNA fragments were separated by agarose ge
electrophoresis. The 5.7 kb fragment corresponding
to P659 without the OCS 3' end was electroeluted from
the gel, extracted with an equal volume of
phenol:chloroform (1:1 v/v) and ethanol
precipitated. This fragment was ligated overnight a
15°C in a volume of 10:1 to the same Bgl
II-blunt/Hind III fragment of P654 used in the
construction of P657. An aliquot of the ligation
mixture was used to transform competent E. coli
JM83. Plasmid DNA from individual ampicillin
resistant colonies was digested with Bam HI and Hind
III until one was found that contained a 4.7 kbp Bam
HI/Hind III fragment. This construction, consisting
of GUS operably linked to a 600 bp P6.1 promoter
fragment and a 1.3 kbp P6.1 3' downstream region
fragment, was designated P660 (Figure 20).

EXAMPLE 12

Construction of Recombinant Genes Under Transcriptional Control of Chimeric Promoters Containing An Inducible Regulatory Element From the Corn 2-2 Promoter

Oligonuclectides were synthesized using an Applied Biosystems Model 380A DNA synthesizer. All oligonuclectides were purified using Applied Biosystems Oligonuclectide Purification Cartridges (cat. # 400771) using the protocol supplied by the manufacturer.

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Construction of pHPH401 and pHPH401 dcm-

Complementary oligonucleotides 32 and 33, of the

- 5 sequences:
 - 32 5'-AATTCGTTAACCGCACCCTCCTTCCCGTCGTTTCCCATCTCTTCCTC CTTTAGA-3'
- 1 n 33 5'-GGAGGAAGAGGTGGGAAACGACGGGAAGGAGGGTGCGGTTAACG-31

and complementary oligonucleotides 34 and 35 of the sequences:

34 5'-GCTACCACTATATAAATCAGGGCTCATTTTCTCGCTCCTCACAGGC CTGGTAC-3'

35 5'-CAGGCCTGTGAGGAGCGAGAAAATGAGCCCTGATTTATATAGTGGT AGCTCTAAA-3'

phosphorylated by incubation of 10 ug of each oligo with 25-50 units of T4 polynucleotide kinase in 50 ul of 50 mM Tris-HCl pH 7.5, 10 mM MgCl2, 10 mM DTT for 20 min at 37°C. An additional 25 ul of 50 mM

25 Tris-HCl pH 7.5, 10 mM MgCl2, 10 mM DTT containing 12.5-25 units of polynucleotide kinase were added and the incubation was continued for 20 min at 37°C.

Kinase reactions were heated to 70°C for 10 minutes and then cooled on ice. Phosphorylated oligos 32,

- 33, 34 and 35 were mixed at a final concentration of 13 ug/ml each in water and 1 ul of this mixture was ligated overnight at 15°C with 1.5 ug of the plasmid vector pBluescript S/K(+) was digested to completion
- 35 with Eco RI and Kpn I and dephosphorylated using calf intestinal alkaline phosphatase. The ligati n mixture was diluted to 60 ul with H2O, and 2 ul of

	the dilution was used to transform 60 ul of competent
	motor cerrs. Aliquots of the transformation roadia-
5	were spread onto LB plates with 50 ug/ml amp and
	plates were incubated overnight at 37°C Small and
	prasmid preparations were performed on amp resistant
	colonies and colonies found to contain 100 be see
	RI/Kpn I insets by restriction digestion were
10	sequenced using the M13 universal primer. One galant
	containing oligonucleotides 32-35 cloned into the Fac
	RI/Kpn site of pBluescript S/K(+) was designated
	PHPH401
	The plasmid pHPH401 was transferred to the dcm-
15	E. Coll Strain NS2616. Any Commonly available de-
	E. COIL strain can be used for this purpose
	Competent NS2216 cells were made by inoculating a so
	mi of LB broth with 100 ul of an overnight auti-
	MOZZIB (grown in LB) and incubating this new gulture
20	at 3/°C with shaking until the Acro reached 0 35
	The culture was chilled to 0°C on ice Rockeria
	marvested by centrifugation at 1500 x g for 10
	minutes, resuspended in 25 ml of 100 mM CaCl- and
	incubated on ice for 30 min. The bacteria were
25	recentrifuged as above and resuspended in 0 5 ml -f
	100 mm CaCl2. After 4 hours on ice. 100 ul of
	competent cells were removed, 4 ng of puphing were
	added, and the cells were incubated on ice for 20
30	minutes. The cells were then heat shocked for 5
30	minutes in a 37°C water bath without shaking. The
	cells were returned to the ice for 2 minutes before
	addition of 2 ml of LB medium. Cells were incubated
	at 3/°C for 1 hour and aliquots of the transformation
35	mixture were plated on LB agar plates containing 50
33	ug/mi amp and allowed to grow overnight at 370 C
	Small scale plasmid preparations from individual amp

	resistant colonies were analyzed by restriction	
5	endonuclease digestions until one was identified that contained pHPH401. The strain was designated HPH401 dcm- and the dcm- plasmid in this strain was designated pHPH401 dcm	
10	Construction of pHPH410 Complementary oligonucleotides 36 and 37 of the sequences:	
15	36 5'-CTCATCAGCACCCCGGCACTGCCACCCCGACTCCCTGCACCTGCCAT GGCTGGGGTCGAGGTAC-3' 37 5'-CTCGAGCCACAGCCATGGCAGGGGTGCAGGGAGTCGGGGTGGCACTGCC GGGGTGCTGATGGCAG GGGGTGCTGATGGCAG	
20	were phosphorylated as above and diluted to 33 ng/ul of each oligo in H_2O . One ul of this dilution was ligated overnight with 1.4 ug Kpn I and Stu I	
25	digested and dephosphorylated pHPH401 dcm- in a volume of 10 ul. The ligation reaction was diluted to 50 ul with water and 2 ul aliquot of the diluted ligation mixture was used to transform 40 ul of competent HB101 cells. Aliquots of the transformation mixture were plated on LB goar plates	
	containing 50 ug/ml amp and allowed to grow overnight at 37°C. Small scale plasmid preparations were prepared from individual amp resistant colonies and colonies found to contain 160 bp Eco RI/Kpn I insets by restriction digestion were sequenced using the M13	:
25	universal primer. One colony containing oligonucleotides 36 and 37 cloned into the Kpn I/Stu	•

35 I sites of pHPH401 dcm- was designated pHPH410.

Construction of the 443 Promoter in pHPH443

Complementary oligonucleotides 44 and 45 of the

- 5 sequences:
 - 44 5'-AATTCTACGTACCATATAGTAAGACTTTGTATATAAGACGTCACC
 TCTTACGTGCATGGTTATATGCGACATGTGCAGTGACGTT-3'
- 10 45 5'-AACGTCACTGCACATGTCGCATATAACCATGCACGTAAGAGGTGA
 CGTCTTATATACAAAGTCTTACTATATGGTACGTAG-3'
- were phosphorylated as above and diluted to 13.3 ng/ul of each oligo in H₂O. One ul of this dilution 15 was ligated overnight with 1.5 ug Hpa I and Eco RI digested and dephosphorylated pHPH410 in a volume of 15 ul. The ligation reaction was diluted to 60 ul with water and 2 ul aliquot of the diluted ligation mixture was used to transform 40 ul of competent
- 20 HB101 cells. Aliquots of the transformation mixture were plated on LB agar plates containing 50 ug/ml emp and allowed to grow overnight at 37°C. Small scale plasmid preparations were prepared from individual amp resistant colonies and colonies found to contain
- 25 240 bp Pst I/Kpn I insets by restriction digestion were sequenced using the M13 universal primer. One colony containing oligonuclectides 44 and 45 ligated into the Eco RI/Hpa I sites of the plasmid pHPH410 was designated pHPH443. The sequence of the insert
- containing in the plasmid pHPH443 is shown in Figure 21. This DNA fragment represents a chimeric promoter consisting of a 77 bp chemically inducible element from the maize 2-2 promoter (nucleotides 9-86 of Figure 21) operably linked to the -1 to -94 of the
- 35 alcohol dehydrogenase 1-18 allele [Dennis et al. (1984) Nucleic Acid Res. 12: 3983-4000] (nucleotides

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87-180 of Figure 21) and using a 5' untranslated region from the corn 2-2 gene (nucleotides 181-225 of Figure 21). The arrow and underlining in the figure

5 denote the transcription and translation start sites. respectively, of the promoter.

Construction of pHPH412

10 Complementary oligonucleotides 46 and 47 of the sequences:

46 5'-CTCATCTCGCTTTGGATCGATTGCTTTCGTAACTGGTGAAGGACTGA

GGCCTAACGGTAC-3' 15

> 47 5'-CGTTAGGCCTCAGTCCTTLACCAGTTACGAAACCAATCGATCCAAAG CGAGATGAG-3'

were phosphorylated as above and diluted to 13.3 20 ng/ul of each oligo in H2O. One ul of this dilution was ligated overnight with 1.4 ug Kpn I and Stu I digested and dephosphorylated pHPH401 dcm- in a volume of 15 ul. The ligation reaction was diluted to 60 ul with water and 2 ul aliquot of the diluted

ligation mixture was used to transform 40 ul of competent HB101 cells. Aliquots of the transformation mixture were plated on LB agar plates containing 50 ug/ml amp and allowed to grow overnight at 37°C. Small scale plasmid preparations were

prepared from individual amp resistant colonies and colonies found to contain 240 bp Kpn I/Pst I I inserts by restriction digestion were sequenced using the M13 universal primer. One colony containing oligonucleotides 45 and 46 cloned into the Kpn I/Stu

35 I sites of the plasmid pHPH401 dcm- was designated PHPH411.

Complementary oligonucleotides 48 and 49 of the sequences: 48 5'-GTCTCGGAGTGGATGATTTGGGATTCTGTTCGAAGATTTGCGGAGG GGGGCCATGGCGACGGTAC-3 49 5'-CGTCGCCATGGCCCCCCCCCGCAAATCTTCGAACAGAATCCCAAAT 10 CATCCACTCCGAGAC-3 were phosphorylated as above and diluted to 20 ng/ul of each oligo in $\rm H_2O$. One ul of this dilution was ligated for 4 hours with 1.4 ug Kpn I and Stu I 15 digested and dephosphorylated pHPH411 in a volume of 10 ul. The ligation reaction was diluted to 50 ul with water and 2 ul aliquot of the diluted ligation mixture was used to transform 40 ul of competent HB101 cells. Aliquots of the transformation mixture 20 were plated on LB agar plates containing 50 ug/ml amp and allowed to grow overnight at 37°C. Small scale plasmid preparations prepared from individual amp resistant colonies were sequenced using the M13 universal primer. One colony containing 25 oligonucleotides 48 and 49 ligated into the Kpn I/Stu I sites of pHPH411 was designated pHPH412. Construction of pHPH460 Complementary oligonucleotides 62 and 63 of the 30 sequences: 62 5'-GTACGTACCATATAGTAAGACTTTGTATATAAGACGTCACCTCTTA CGTGCATGGTTAACA-3

53 5'-AGCTTGTTAACCATGCACGTAAGAGGTGACGTCTTATATACAAAGT CTTACTATATGGTACGTACTGCA-3'

5	were phosphorylated as above and mixed together at 10 $\rm ng/u1$ of each oligo in $\rm H_2O$. One u1 of this dilution	
	was ligated for 6 hours with 1 ug Pst I and Hind III	
	digested and dephosphorylated pBluescript S/K(+) in a volume of 10 ul. The ligation reaction was diluted	Ţ,
10	to 50 ul with water and 2 ul aliguot of the diluted ligation mixture was used to transform 40 ul of competent HB101 cells. Aliguots of the	
	transformation mixture were plated on LB agar plates containing 50 ug/ml amp and allowed to grow overnight	
15	at 37°C. Small scale plasmid preparations made from individual amp resistant colonies were sequenced	
	using the M13 universal primer until a colony containing oligonucleotides 62 and 63 cloned into the Pst I/Hind III sites of pBluescript S/K(+) was	
20	found. This plasmid was designated pHPH460	

Construction of pHPH461

Complementary oligonucleotides 75 and 76 of the sequences:

25 75 5'-ATATGCGACATGTGCAGTGACGTTATCAGATATAGCTCACCCTATAT ATATAGCTCTGTCCGGTGTCGAC-3'

76 5'-TCGAGTCGACACCGGACAGAGCTATATATATAGGGTGAGCTATATCT GATAACGTCACTGCACATGTCGCATAT-3'

30

were phosphorylated as above and mixed together at 12.5 ng/ul of each oligo in H₂O. One ul of this dilution was ligated for 6 hours with 1 ug Hpa I and Xho I digested and dephosphorylated PHPH460 in a volume of 10 ul. The ligation reaction was diluted to 50 ul with water and 2 ul aliquot of the diluted

ligation mixture was used to transform 40 ul of competent HB101 cells. Aliquots of the

- transformation mixture were plated on LB agar plates containing 50 ug/ml amp and allowed to grow overnight at 37°C. Small scale plasmid preparations made from individual amp resistant colonies were sequenced using the M13 universal primer. One colony
- 10 containing oligonucleotides 75 and 76 cloned into the Hpa I/Xho I sites of pHPH460 was designated pHPH461.

Construction of pHPH462

Complementary oligonucleotides 77 and 78 of the sequences:

77 5'-AAGTGACAATCACCATTCATCTCGCTTTGGATCGATTGGTTTCGTAA CTGGTGAAGGACTGAGGCCTAACGGTAC-3'

20 78 5'-CGTTAGGCCTCAGTCCTTCACCAGTTACGAAACCAATCGATCCAAAC GAGATGAATGGTGATTCTCACT-3'

were phosphorylated as above and mixed together at 10 ng/ul of each oligo in H₂O. One ul of this dilution 19/ul of each oligo in H₂O. One ul of this dilution was ligated for 6 hours with l ug Kpn I and Hinc II digested and dephosphorylated pHPH461 in a volume of 10 ul. The ligation reaction was diluted to 50 ul with water and 2 ul aliquot of the diluted ligation mixture was used to transform 40 ul of competent 40 lblook 10 lbloo

35 universal primer. One col my containing

oligonucleotides 77 and 78 cloned into the Kpn I/Hinc II sites was designated pHPH462

5

10

15

Construction of pHPH463 and pHPH463dam-

Phosphorylated, complementary oligonucleotides 48 and 49, described above, were mixed together at 25 mg/ul of each oligo in H₂O. One ul of this dilution was ligated for 6 hours with 1 ug Stu I and Kpn I digested and dephosphorylated pHPH462 in a volume of 10 ul. The ligation reaction was diluted to 50 ul with water and 2 ul aliquot of the diluted ligation mixture was used to transform 40 ul of competent HB101 cells. Aliquots of the transformation mixture were spread on LB agar plates containing 50 ug/ml amp and allowed to grow overnight at 37°C. Small scale

plasmid preparations made from individual amp
resistant colonies were sequenced using the M13

universal primer. One colony containing oligonucleotides 48 and 49 cloned into the Stu I/Kpn I sites of pHPH462 was designated pHPH463. The sequence of the insert contained in the plasmid pHPH463 is shown in Figure 22. This DNA fragment represents a chimeric promoter consisting of the -1

25 represents a chimeric promoter consisting of the -1 to -136 region of the 2-2 promoter (nucleotides 7-146 of Figure 22) operably linked to the 5' untranslated leader from the maize alcohol dehydrogenase 1-15 allele [Dennis et al. (1984) Nucleic Acids Res. 12:

3983-4000] (nucleotides 147-247 of Figure 22) and modified to incorporate an Nco I site at the translation state codon. The arrow and underlining in the figure denote the transcription and translation start sites, respectively, of the

35 promoter.

	Ti	ıe	plasm	iđ p	нрн4 (3 was	transform	ned	into	the
dam-	E.	CC	li st	rain	CHS	26-usi	g the pro	Cec	lure	
desc	ribe	đ	above	for	the	transi	ormation	of	пНРН	101

described above for the transformation of pHPH401 into the dcm- E. coli strain NS2216. The plasmid pHPH463 in E. coli CHS26 was designated pHPH463dam-.

Construction of pHPH467

-3.

10 Complementary oligonucleotides 88 and 89 of the sequences:

88 5'-GTTANCANGGATCGGCGCCCCCCCCCGCGGCTCGCCCCTATATTTATA TTTGCTCANTGGACAGGCATGGGGCCTATCTCGCTTTGGAT-3'

89 5'-CGATCCAAAGCGAGATAGCCCCATGCCTGTCCATTGAGCAAATATA
AATATAGCGGCGAGCTCGGCGGGGGGGGGCGCGATCCTTGTTAACTGCA

- 20 were prepared and phosphorylated by incubation of 5 ug of each oligo with 25-50 units of T₄ polynucleotide kinase in 50 ul of 50 mM Tris-HCl pH 7.5, 10 mM MgCl₂, 10 mM DTT for 1 hour at 37°C. Kinase reactions were heated to 70°C for 10 minutes
- and then cooled on ice. Oligonucleotides were mixed together at 35 ng/ul of each oligo in H₂O. One ul of this dilution was ligated for 4 hours with 1 ug of Pst I and Cla I digested and dephosphorylated PHPH463dam— in a volume of 10 ul. The ligation
- 30 reaction was diluted to 50 ul with water and 2 ul aliquot of the diluted ligation mixture was used to transform 40 ul of competent HBI01 cells. Aliquots of the transformation mixture were plated on LB agar plates containing 50 ug/ml amp and allowed to grow
- 35 overnight at 37°C. Small scale plasmid preparations made from individual amp resistant colonies were

the Bam HI site of the vector polylinker was designated pHPH500.

Construction of pHPH478

The plasmid pHPH500 was digested to completion with Bam HI and Hpa I. The digestion products were separated by polyacrylamide gel electrophoresis and the 85 bp fragment corresponding to the inducible

- the 85 bp fragment corresponding to the inducible element of the 2-2 promoter was recovered as described above. This fragment was ligated overnight with 1 ug of Bam HI and Hpa I digested and dephosphorylated pHPH467 in a volume of 10 ul. The
 - 5 ligation reaction was diluted to 50 ul with water and 2 ul aliquot of the diluted ligation mixture was used to transform 40 ul of competent HBIOI cells. Aliquots of the transformation mixture were plated on
- LR agar plates containing 50 ug/ml amp and allowed to grow overnight at 37°C. Small scale plasmid preparations made from individual amp resistant colonies were analyzed by restriction endonuclease digestion until a plasmid containing the Bam HI/Hpa I fragment of pHPHS00 cloned into the Bam HI/Hpa I
- 25 sites of pHPH467 was identified. This plasmid was designated pHPH478. The sequence of the insert contained in the plasmid pHPH478 is shown in Figure 23. This DNA fragment represents a chimeric promoter consisting of a 76 bp chemically inducible element
- of Figure 23) operably linked to the -1 to -94 region of the phytochrome type 3 promoter (Hershey et al. (1987) Gene 61: 339-348] (nucleotides 86-155 of Figure 23) and using a 5' untranslated region from
- 35 the maize alcohol dehydrogenase 1-15 allele [Dennis et al. (1984) Nucleic Acids Res. 12: 3983-4000]

(nucleotides 156-256 of Figure 23) and modified to incorporate an Neo I site at the translation start

5 codon. The arrow and underlining in the figure denote the transcription and translation start sites, respectively, of the promoter.

Construction of pHPH443GUS, pHPH410GUS, pHPH412GUS, pHPH412GUS, pHPH463GUS and pHPH478GUS

Sixty micrograms of the plasmid pHPH443 were digested to completion with Xba I and Nco I. The resulting DNA fragments were separated by

- electrophoresis overnight at 180 V in a single 1 cm wide lane of a 2 mm thick 7.5% polyacrylamide gel made in TBE and containing 25% glycerol. DNA fragments were visualized under UV light after staining the gel in 0.5 ug/ml ethidium bromide in H₂O. for 20 minutes. The 230 bp DNA fragment
- corresponding to the insert of pHPH443 was excised from the gel with a scalpel, placed in a 1.5 ml microcentrifuge tube, crushed with a spatula and suspended in gel elution buffer. The tube was then shaken vigorously overnight at 37°C. Gel fragments were removed from the resulting slurry by finishing
- 25 were removed from the resulting slurry by filtration through glass wool and DNA in the filtrate was precipitated on dry ice after adding 1 ml of ethanol. DNA was recovered by centrifugation and resuspended by vigorous vortexing in 0.3 ml of TE pH 30 8.0. The suspension was centrifuged and the
 - 8.0. The suspension was centrifuged and the supernatant was transferred to a new tube, made 0.3 M in sodium acetate and precipitated on dry ice as described above. DNA was collected by centrifugation, and the pellet was dissolved in 20 ul
- 35 of TE pH 7.5 after being dried in vacou. A 0.5 ul aliquot of pHPH443 insert DNA were ligated to 1 ug

15

- Xba I and Nco I digested and dephosphorylated pTD136 (Example 8) in a volume of 10 ul. The ligation reaction was diluted to 50 ul with water and 2 ul aliquot of the diluted ligation mixture was used to transform 40 ul of competent HB101 cells. Aliquots of the transformation mixture were plated on LB agar plates containing 50 ug/ml amp and allowed to grow overnight at 37°C. Small scale plasmid preparations were performed on amp-resistant colonies and the resulting DNAs were digested with Xba I and Nco I until a colony was found that contained the 230 bp Xba I/Nco I fragment from pHPH443 in pTD136. This plasmid, consisting of the promoter fragment of pHPH443 operably linked to the GUS/2-1 3' end construction in pTDS136 was called pHPH443GUS. Similarly, the Xba I/Nco I promoter fragments
- of pHPH410. pHPH412, pHPH463 and pHPH478 were cloned into the Xba I/Nco I sites of pTDS136 to create the plasmids pHPH410GUS, pHPH412GUS, pHPH463GUS and pHPH478GUS. respectively.

Construction of pHPH420GUS

- Thirty micrograms of the plasmid pHPH412 were digested to completion with Hpa I and Nco I. The resulting DNA fragments were separated by electrophoresis overnight at 250 V in a single 1 cm wide lane of a 2 mm thick 7.5% polyacrylamide gel made in TBE and containing 25% glycerol. DNA fragments were visualized under UV light after staining the gel in 0.5 ug/ml ethidium bromide in H₂O for 20 minutes. The 200 bp DNA fragment corresponding to the insert of pHPH412 was recovered
- 35 from the gel as described above and dissolved in 20 ul of TE pH 8.0 The concentration of the pHPH412

	40 ng of pHPH412 insert DNA were ligated to 1 ug Hpa
5	I and Nco I digested and dephosphorylated pHPH443GUS
	in a volume of 10 ul. The ligation reaction was
	diluted to 50 ul with water and 2 ul aliquot of the
	diluted ligation mixture was used to transform 40 ul
	of competent HB101 cells. Aliquots of the
10	transformation mixture were plated on LB agar plates
	containing 50 ug/ml amp and allowed to grow overnight
	at 37°C. Small scale plasmid preparations were
	performed on amp-resistant colonies and the resulting
	DNAs were digested with Xba I and Nco I until a
15	
	colony was found that contained the 200 bp Hpa I/Nco
	I promoter fragment from pHPH412 in pHPH443 GUS.
	This plasmid construction was called pHPH420GUS. The
 	sequence of the insert contained in the plasmid
	pHPH420 is shown in Figure 24. This DNA fragment
20	represents a chimeric promoter consisting of a 77 bp
	chemically inducible element derived from the maize
	2-2 promoter (nucleotides 9-86 of Figure 24) operably
	linked to the -94 to +101 region of the maize alcohol
	dehydrogenase 1-1S allele [Dennis et al. (1984)
25	Nucleic Acids Res. 12: 3983-4000] (nucleotides 87-281
	of Figure 24) and modified to incorporate an Nco I
	site at the translation start codon. The arrow and
	underlining in the figure denote the transcription
	and translation start sites, respectively, of the

30 promoter.

EXAMPLE 13

Construction of Recombinant Promoters Containing Various Modifications of the 2-2 Inducible Element

Construction of plasmids pAl-pA70

Individual oligonucleotides incorporating various base changes at one or more positions in their sequences were prepared using an Applied Biosystems Model 380A DNA synthesizer by using mixtures of nucleoside phosphoramidites at specific cycles in the synthesis. In a similar manner, populations of complementary oligonucleotides to

15 those made above were prepared by incorporating mixture of nucleoside phosphoramidites at appropriate synthesis cycles so as to complement the possible base heterogeneities in the first strand.

- 20 The complementary pairs of oligos-:
 - 103 5'-CACCTCTTACGTGCATGGTTANATGNNACATNIGCAGTGANGTT-3'
 104 5'-AACNICACTGCANATGINNCAINTAACCATGCACGTAAGAGGTGA
- 25
 - 105 5'-CACCTCTTACGTGCATGGTTAIATGCGACATGTGNAGTPACGTT
 106 5'-AACGTRACTNCACATGTCGCATATAACCATGCACGTAAGAGGTG
 ACGT-3'
- 30 107 5'-CACCICTIACGIGCATGGITATATGCGACARGIGCPPRGACGIT
 108 5'-AACGICPRRGCACPTGICGCATATAACCATGCACGTAAGAGGIG
 ACGI-3'
- 109 5'-CACCTCTTACGTGCATGGTTATATGCGPRPTGTGCAGTGACGTT
 35 110 5'-AACGTCACTGCACGRPRCGCATATAACCATGCACGTAAGAGGTG
 ACGT-3'

 	111 5'-CACCTCTTACGTGCATGGTTATATGCGACATGRPCAGTGPCGTT	
5	112 5'-AACGRCACTGRPCATGTCGCATATAACCATGCACGTAAGAGGTG	
	ACGT~3'	:
	115 5'-CACCTCTTACGTGCATGGTTPTPRPCGACATGTGCAGTGACGTT	•
	116 5'-AACGTCACTGCACATGTCGRPRARAACCATGCACGTAAGAGGTG	
10	ACGT-3'	
	117 5'-CACCTCTTACGTGRARGPTRATATGCGACATGTGCAGTGACGTT	
	118 5'-AACGTCACTGCACATGTCGCATATPARCPTPCACGTAAGAGGTG	
	ACGT-3'	
15		
	where	
	N= A,C,G,T	
	P=A,G	
	R=C,T	
20		
	were phosphorylated as described in Example 12 and	
	each pair was ligated in equimolar ratios with	
	pHPH443GUS that had been digested to completion with	
	both Hpa I and Aat II and dephosphorylated. The	
25	ligation reactions were diluted to 50 ul with water	
	and 2 ul aliquot of the diluted ligation mixtures	
	were used to transform 40 ul of competent HB101	
	cells. Aliquots of the transformation mixtures were	
	plated on LB agar plates containing 50 ug/ml amp and	
30	allowed to grow overnight at 37°C. Small scale	
	plasmid preparations were need	
	plasmid preparations were performed on amp-resistant	
	colonies and the resulting DNAs were sequenced using	,
	either oligo 35 (Example) or oligo HH114 primer	•
35	(HH114 sequence: 5'-GGAGGAAGAGATGGGAAACGACGGG-3').	•
	Plasmids in which base changes had been introduced in	•
	the region of PHPH443GUS corresponding to the 77 bp	

		inducible element from the 2-2 promoter were
		selected. Table lists the plasmids that
	5	single base changes in the region of interest.
•		Similarly, the complementary pairs of
:		oligonucleotides-:
		121 5'-CTAGTGAATTCGTACCATATAGRAAGPCRRTGTATATAAGACGT-3'
	10	122 5'-CTTATATACAPPGRCTTPCTATATGGTACGAATTCA-3'
		123 5'-CTAGTGAATTCGTACCATATAGTAAGACTTRPRATPTAAGACGT-3'
		124 5'-CTTARATPRPAAGTCTTACTATATGGTACGAATTCA-3'
	15	
	13	125 5'-CTAGTGAATTCGTACCATATAGTAAGACTTTGTPRATPPGACGT-3'
		126 5'-CRRATRPACAAAGTCTTACTATATGGTACGAATTCA-3'
		127 EL CR
		127 5'-CTAGTGAATTCGTACCATARAPTPAPACTTTGTATATAAGACGT-3'
	20	128 5'-CTTATATACAAAGTETRARTPTATGGTACGAATTCA-3'
		Where:
		N= A,C,G,T
		P=A,G
		R=C,T
	25	
		were phoshphorylated as described in Example and
		rigated in equimolar ratios with puppers cur
		seen digested with both Xba I and Aa+ II and
		dephosphorylated. The ligation reactions
	30	transformed into HBIO1 cells and plasmids in the
		base changes had been introduced in the region of
		phra443GUS corresponding to the 77 hp industria
•		element from the 2-2 promoter were solvents
•		described above. Table 3 lists the plasmids that
	35	were round to contain base changes in the 77 bp
		industry of

inducible element from the 2-2 promoter listed below and the p sitions of those those changes.

TGTACACGTCACTGCAA

5	1	GIACCATATAGTAAGACTTIGTATATAAGACGTCACCTCTTACGTGCATGGTTATATGCG	60	,
		CATGGTATATCATTCTGAAACATATATTCTGCAGTGGAGAATGCACGTACCAATATACGC		•
		ACATGTGCAGTGACGTT		•

Table 3

15		Nucleot	ide chang	•	
	Plasmid pA #	_F.com_	-	Position	
	0	No	changes		
	1	G	С	70	
20	2	Ŧ	С	64	
		G	С	70	
		T	A	71	
	3	A	G	69	
	4	A	G	69	
25		G	A	70	
		T	С	71	
	5	T	С	64	
		A	G	69	
		T	С	71	
30	6	T	С	64	
	7	T	A	55	
		С	T	59	
		G	A	60	•
		G	A	65	
35		c	A	74	•

		8	T	T	55	
			G	т-	60	
:	5		С	A	74	
*		9	С	т	74	
:		10	· т	G	55	
•			С	A	55 59	
			С	G	74	
	10	11	c	G	74 59	
			G	T		
			G	T	60	
		12	c	G	74	
			G	A	59	
178	15	13	T	A	65	
			c		55	
			G	G	59	
			. с	A	65	
		14	Ţ	T	74	
	20		C T	С	55	
				Ŧ	59	
		15	G	T	65	
			c	A	68	
		16	G	A	72	
	25		С	T	68	
		17	G	A	72	
		1/	С	G	68	
		18	G	A	72	
			С	G	68	
	30	19	С	T	68	•
	30	20	A	G	61	
			С	T	62	
;			A	G	63	
•		21	A	G	61	
:		22	. с	T	62	
-	35	23	A	G	61	
			A	G	63	
					• • • • • • • • • • • • • • • • • • • •	

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			16	6	PCT/US90/01210	
	24		A	G	63	
5	25		С	G	62	
3			A	T	63	
	26		G	A	67	:
	27		T	С	66	
			G	A	67	:
	28		T	С	66	
10			G	A	67	
			A	G	73	
	29		A	G	73	
	31		T	С	66	
			Ą	G	73	
15	32		A	G	67	
			A	G	73	
	33		T	С	66	
	34	•	A	G	54	
			T	С	57	
20	35		A	G	16	
			T	С	18	
	36		A	G	16	
	37		T	G	19	
	38		T	С	18	
25	39		T	С	18	
			T	T	19	
	40		T	С	12	
	41		G	A	21	
			T	С	22	
30			A	G	25	
	42		G	A	21	
	43		T	С	20	
			G	A	. 21	:
	44		T	C	20	
35	45		Ŧ	c	12	:
			A	G	16	
				-	10	

С

18

-0.4			10	57	PCT/US90/01210	
		46 47	T A	c	22	
	5_	48	G	G G	25	
:			T	T	21	
		49	T	C	22	
;				С	20	
			T	С	22	
	10	50	A -	G	25	
			T	С	20	
			G	A	21	
		51	T	С	22	
			A	T	56	
	15	52	T	С	57	
-		53	G	A	58	
		54	A	G	56	
		55	A	G	54	
		33	. A	G	54	
	20	56	A	- G	56	
		57	T	С	12	
		58	G	С	25	
		59	T	С	24	
		• ,	A	G	23	
	25	60	T	С	24	
		00	A	G	23	
		61	A	G	28	
		62	A	G	23	
		02	A	G	27	
	30	63	A	G	28	
		64	T	С	9	
		65	G	A	11	
:		65	G	A	11	
•			G	A	15	
	35	66	T	С	9	
	33		G	A	11	
		67	A	G	13	
					==	

		168			
	68	G	A	11	
		A	G	13	
5	69	G	A	15	
	70	С	T	47	:

EXAMPLE 14

The Use of N-(aminocarbonyl)-2-chlorobenzene-

sulfonamide to Induce the Expression of Recombinant GUS/2-1 Corn Gene Constructions in Transformed Rice Protoplasts

Transformation of Rice Protoplasts

- Rice suspension cultures, initiated from anther-derived callus, were maintained by weekly subculture at a 1:4 dilution ratio with fresh liquid N6 medium as described by Chu et al. [Sci Sinice 18:659-668 (1975)] containing 2 mg/ml 2,4-dichloro-
- phenoxyacetic acid and 3% (w/v) sucrose, pH 6.0.
 Protoplasts were isolated from suspensions of rice
 cells 4-6 days after subculture by overnight
 incubation (16-18 hrs) in 4 ml of enzyme solution (2%
 (w/v) cellulose "Onozuka" RS and 0.5% (w/v)
- 25 Macerozyme (both from Yakult Honsha, Nishinomiya, Japan), 13% (w/v) mannitol, pH 5.6) per gram of cells and agitation of the mixture on a rotary shaker at 30 rpm at 25 C. Released protoplasts were filtered through a 60 mm mesh size nylon screen, transferred
- 30 to 50 ml Pyrex® test tubes and washed twice by centrifugation at 80 g for 10 minutes in Kren's F solution (140 mM NaCl), 3.6 mM KCl, 0.75 mM·Na₂HPO₄ 7H₂O, 5 mM glucose, 125 mM CaCl₂, pH 7.0).
- Protoplasts were purified by resuspending the pellet 35 in N6 medium with 17% (w/v) sucrose, centrifuging at 80g for 20 minutes and collecting the floating

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layer. Cell counts were made with a Fuchs-Rosenthal hemocytometer.

Protoplasts were transformed as follows:

Multiple aliquots of the protoplasts (5-10 x 106 cells) were centrifuged gently (80 g) for 4 minutes in sterile tubes. The supernatant was discarded and the cells were resuspended in 1 ml of Kern's F, pH 5.8 buffer. Ten μg of transforming DNA in less than 15 μl of TE pH 8.0 were added per million protoplasts. The tubes were shaken gently to disperse the cells in the DNA solution, and 0.6 ml of a solution containing 40 % PEG (Polysciences Inc., Warrington PA 18976, CAT # 1102) and 3 mM CaCl2 was

Warrington PA 18976, CAT # 1102) and 3 mM CaCl₂ was added. The resulting protoplast cell suspension was mixed gently and incubated at room temperature for 20 minutes. A volume of 13-15 mls of Kren's F, pH 7.0 solution was then added to dilute out PEG.

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N-(aminocarbonyl)-2-chlorobenzenesulfonamide Induction of Transformed Rice

The transformed protoplasts were collected by centrifugation at 80g for 4 minutes. The supernatant 25 was discarded and the protoplasts were resuspended in 2.0 ml of Kren's F, pH 5.8. The protoplast sample was divided into two 1 ml aliquots. One ml of protoplast medium was added to one aliquot of the protoplasts, while 1 ml of the protoplast medium containing 100 µg/ml N-(aminocarbony1)-2-chlorobenzenesulfonamide was added to the other aliquot. Protoplasts were then incubated at 25°C in the dark for 16 hours.

The inducibility of the recombinant GUS genes whose expression were controlled by 2-1 corn gene promoter and downstream sequences were determined by

	measuring the level of the β-glucuronidase enzyme	
5	activity in protoplasts cultured in the presence and absence of N-(aminocarbonyl)-2-chlorobenzene.	
	sulfonamide. GUS activity was assayed by harvesting	•
	protoplasts in a clinical centrifuge at 80g for 5	•
	minutes, and resuspending them in 1.0 ml 1X GUS lysis	•
10	buffer (50 mM sodium phosphate pH 7.0, 10 mM	
	β-mercaptoethanol, 10 mM EDTA, 0.1% Triton X-100,	
	0.1% N-lauroylsarcosine). The suspension containing the lysed protoplasts was vortexed and spun at top	
	speed in a table top clinical centrifuge for 5	
	minutes. Eighteen µl of the supernatant was	
15	transferred to a tube containing 782 µl of water for	
	determination of protein content in the protoplast	
	lysate. Protein content in the diluted lysate was	
	determined using the Bio-Rad Protein Assay kit	
	(Blo-Rad Laboratories, Richmond, CA 94804) following	
20	the manufacturer's recommendations for the migrogram	
	procedure. A protein concentration curve was	
	prepared using bovine serum albumin as a standard	
	The protein content, so determined, was multiplied by	
	a factor of 7.2 to give the protein content in 120	
25	or extract (the amount of extract present in a simple	
	time point of the assay-see below). Of the remaining	

supernatant, 585 µl was transferred to a fresh tube.

The substrate for the GUS assay was
4-methyl-umbelliferyl-D-D-glucuronide (4-MUG) and was
Obtained from Sigma Chemical Co., St. Louis MO 63178
(CAT # 9130). 4-MUG was prepared as a 10 mM stock in

IX GUS buffer. Sixty-five µl of a pre-warmed (37°C)
10 mM 4-MUG stock was added to the pre-warmed 585 µl
protoplast extract, and a 100 µl aliquot of the
resulting mixture was transferred to a well of a
24-well microtiter dish containing 0.9 ml of 0.2 M

		Na ₂ CO3. Similar aliquots are removed at 1 hour, 2
:	5	hour, and 3 hours. The 4-MU flowrescence of individual samples, from each time point was determined quantitatively using an excitation wavelength of 365 nm and measuring flourescence at an emission wavelength.
•	10	emission wavelength of 455 nm. A standard curve of 4-MU flourescence was also prepared by measuring the flourescence of 100 nM and 1 uM 4-MU (Sigma Chemical Co., CAT # 1508). GUS activity in the transient assay was expressed as picomoles of 4-MU produced per up protein per hour.
	15	The results of transient assays of the type described above are summarized in Table 4 for the plasmid constructions pJE 516, pDuPE2, pDuPE8, pDuP19, pDuP19 and pDup113. Plasmid pBM117 was also run in each assay as a control for convenience.
	20	expression. The plasmid consists of a GUS coding region under the control of CaMV 35S promoter and 3 downstream regions.

- Tegion under the control of CaMV 35S promoter and 3 downstream regions. GUS activity resulting from transcription driven by the 2-1 promoter and downstream regions (puB516) was consistently highly induced by addition of 100 µg/ml of
- N-(aminocarbonyl)-2-chlorobenzenesulfonamide to the protoplast medium.

TABLE 4

10	Promoter Sample Size (in bp)	GUS ACTI (FU/µg-n Uninduced* ND 241.1 569.4 227.3 121.1 106.6	nin.)	Fold Induction 0 x 2.6 x 7.6 x 7.9 x 5.9 x 0 x	:
15	*Induction in table	was a a.			
٠	benzenesulfonamide to tr	N-(aminocart		e nloro-	
20	The Use of N-(aminocarbon sulfonamide to Induce Ext Corn Promoter/GUS Gene Co Rice Protoplasts	nyl)-2-chloro		2-2 Imed	
25	Rice suspension cul anther-derived callus, we protoplasts for the trans expression assays. The m transformation and chemic as well as GUS	re utilized ient transfo	as the sou	đ	
30	as well as GUS assays wer Protoplasts were transform 2-2 promoter/GUS fusions of The induction of pTM pDuPM17, pDuPN27, pDuPM4 a CORSTURIEDS (CORSTURIEDS)	e described in med with pBM described belocated by the belocated belocated by the belocated belocated by the belocated belocated by the belocated by th	in Example 1117, and t low.	14. the	
35	constructions (all describ N-(aminocarbony1)-2-chloro transfromed protoplasts we transient expression assay	Ded in Exampl Obenzenesulfo	namide in		:

N-(aminocarbonyl)-2-	-chlorobenzenesulfonamide
inducibility of our	remesultonamide
transformed	expression in protoplasts

- 5 transformed with these constructions is presented in Table 5. The results show that the chemical strongly induces expression of all constructions with 2-2 promoter fragments that are longer than 208 bp. A rapid loss of chemical inducible GUS activity occurs 10
- when the size of the 2-2 promoter fragment is less than 208 bp 5' to the translation start site in the promoter. This indicates that there is a DNA element in the 2-2 promoter contained, at least in part, between nucleotides -210 and -130 bp of 5' of the
- 15 translation start site of the GUS gene that appears necessary for induction of 2-2 promoter activity by $N\hbox{-(aminocarbony1).-2-chlorobenzene sulfonamide.}\\$

20			TABLE 5		
25	Sample NO DNA pBM117 pTDS130 pTDS133 pTDS134 pDUPM17 pDUPN27	Promoter <u>Size (in hp</u>) N/A N/A ~1900 465 450 248 208	GUS ACTI (FU/µg- Uninduced* ND 1.68 1.38 1.52 1.65 1.25		Fold Induction 0.0 x 4.5 x 64.1 x 67.7 x 47.4 x 80.5 x 82.8 x
	pDuPN7	130	0.83 0.54	24.3 1.52	29.0 x 2.8 x

*Induction in Table 5 was accomplished by the addition of 100 µg/ml of N-(aminocarbonyl)-2-chloro-

35 benzenesulfonamide to transformed prot plasts.

EXAMPLE 16

The Use of N-(aminocarbonyl)-2-chlorobenzene
sulfonamide to Induce Expression of Recombinant 5-2

Corn Promoter/GUS Gene Constructions in Transformed
Rice Protoplasts

Rice suspension cultures, initiated from anther-derived callus, were utilized as the source of protoplasts for the transient transformation and expression assays. The method for isolation and transformation of protoplasts, and the GUS assays were described in Example 14. Protoplasts were transformed with pBM117 and the 5-2 promoter/GUS fusions described helow.

The response of pMC 715.53 was analyzed by transient expression assay in rice protoplasts. No induction of GUS expression was observed in

transformed protoplasts treated with N-(aminocarbonyl)-2-chlorobenzenesulfonamide. Since the in vivo induction in the 5-2 gene is the weakest of all corn genes tested, it may be possible that its inducibility cannot be measured in a transient assay.

EXAMPLE 17

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The Use of N-(aminocarbonyl)-2-chlorobenzenesulfonamide to Induce A Chimeric 218 Corn Promoter/GUS Fusion in Transformed Rice Protoplasts

Rice suspension cultures, initiated from anther-derived callus, were utilized as the source of protoplasts for the transient transformation and expression assays. The method for isolation, transformation and chemical treatment of protoplasts, as well as GUS assays are described in Example 14.

The induction	of GUS	activity	in	response	to
treatment of rice pr	otopla	sts trans:	For	ned with	

5 pTDS130 (Example 8) and pMC7113 with 100 mg/l ofN-(aminocarbonyl)-2-chlorobenzenesulfonamide was analyzed by transient expression. The results are presented in Table 6.

10

TABLE 6

	Conchus	GUS ACT	*YTIVI	FOLD
	Construction	UNINDUCED	INDUCED	INDUCTION
15	No DNA	3	3	0
	PHPH130	17.3	546	31.5
	pMC7113	78.4	952	12.1

* GUS activity expressed as flourescence units/hr/106protoplasts

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The results in Table 6 show that GUS activity resulting from transcription of a GUS gene under the control of the the 218 promoter was consistently induced by addition of 100 ug/ml of N-(aminocarbony1)-2-chlorobenzenesulfonamide to the protoplast medium

EXAMPLE 18

The Use of N-(aminocarbonyl)-2-chlorobenzenesulfonamide to Induce Expression of Recombinant P6 Petunia Promoter/GUS Gene Constructions in Transformed Rice Protoplasts

Rice suspension cultures, initiated from anther-derived callus, were utilized as the source of pr t plasts for the transient transformation and expression assays. The method for isolation and

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transformation of protoplasts, and the GUS assays		
were described in Example 14. Protoplasts were		
transformed with pBM117 and various P6.1		
promoter/GUS fusions described below.	:	
The responses of P655, P657, P658, and P660	_	
were analyzed by transient expression assay in rice	· ·	
protoplasts. The degree of induction of GUS		
expression in transformed protoplasts in response to		
N-(aminocarbonyl)-2-chlorobenzenesulfonamide		
treatment is presented in Table 7. GUS activity		
resulting from transcription driven by the P6		
promoter and various 3' downstream regions was		
consistently induced by addition of 100 µg/ml of		
N-(aminocarbony1)-2-chlorobenzenesulfonamide to the		
protoplast medium.		

In addition, all DNA sequences required for this induction appear to reside in the P6.1 promoter,

20 since substitution of a 3' end from a non-inducible gene (the OCS gene) had no effect on the induction of the P6.1 promoter/GUS construction.

TABLE 7

25	Sample	P6 Promoter Size (in bp)	GUS ACTI (FU/µg-m Uninduced*	Fold Induction	
30	NO DNA pBM117 P655 P657 P658 P660	N/A N/A 1300 1300 300 600	ND 174.8 61.8 66.6 64.5 112.8	27.0 325.7 317.6 488.1 404.3	0.0 x 1.9 x 5.1 x 7.3 x 6.3 x

*Induction in Table 7 was accomplished by the addition of 100 ug/ml of N-(aminocarbonyl)-2-chlorobenzenesulfonamide to transformed protoplast.

35 ND= not determined

EXAMPLE 19

The Use of N-(aminocarbonyl)-2-chlorobenzenesulfonamide to Induce Recombinant Genes Under Transcriptional Control of Chimeric Promoters Containing An Inducible Element From the Corn 2-2 Promoter in Transformed Rice Protoplasts

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Rice suspension cultures, initiated from anther-derived callus, were utilized as the source of protoplasts for the transient transformation and expression assays. The method for isolation.

15 transformation and chemical treatment of protoplasts, as well as GUS assays were described in Example 14.

The induction of GUS activity in response to treatment of rice protoplasts transformed with pTDS130, pHPH410GUS, pHPH412GUS, pHPH42GUS,

TABLE 8

20 PHPH443GUS, PHPH463GUS and PHPH478GUS with N-(aminocarbonyl)-2-chlorobenzenesulfonamide was analyzed by transient expression. The results are presented in Table B.

25 GUS ACTIVITY* FOI.D CONSTRUCTION UNINDUCED INDUCED INDUCTION No DNA 10 n pTDS130 124.4 849.2 6.8 pHPH410GUS 17 14 . R pHPH412GUS 21.5 30 26.6 1.2 pHPH420GUS 317.5 1674.5 5.3 pHPH443GUS 55.0 590.6 10.7

90.7

13

35

pHPH463GUS

pHPH478GUS

781.3

160.7

8.6

12.4

^{*} GUS activity expressed as flourescence units/hr/10 6 protoplasts

These results demonstrate that addition of the

5 77 bp element derived from the corn 2-2 promoter to
the promoters regions of non-inducible GUS genes
causes these gene to dispaly induciblity when assayed
in transformed rice protoplasts treated with 100
ug/ml of N-(aminocarbonyl)-2-chlorobenzenesulfonamide.

EXAMPLE 20

The Use of N-(aminocarbonyl)-2-chlorobenzenesulfonamide to Induce Recombinant Genes under the Transcriptional Control of Recombinant Promoters Containing Various Modifications of the 77 bp 2-2 Inducible Element in Transformed Rice Protoplasts

Rice suspension cultures, initiated from
anther-derived callus, were utilized as the source of
protoplasts for the transient transformation and
expression assays. The method for isolation,
transformation and chemical treatment of protoplasts,
as well as GUS assays were described in Example 14.

The induction of GUS activity in response to N-(aminocarbony))-2-chlorobenzenesulfonamide treatment of rice protoplasts transformed with pAO- pA70 was analyzed by transient expression. The degree of induction of GUS expression in transformed

protoplasts in response to N-(aminocarbonyl)-chlorobenzenesulfonamide treatment is presented in Table 9.

25

30

Table 9

PA PROMOTER UNINDUCED* INDUCED* INDUCTION		5		GUS ACTI	VITY	FOLD
No DNA 10 8 0 PHPH443 89 954 10.7 PAO 115 1100 9.5 10 1 86 1001 11.6 2 25.5 72 2.8 3 29.5 495 16.8 4 22 70.5 3.2 5 39 111.5 2.9 15 6 20 106 5.3 7 20 37.5 1.9 8 15 31.5 2.1 9 75 1465 19.5 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 18 32 1.8 14 317 2280 7.2 15 18 32 1.8 17 8.5 23.5 2.8 18 19 161 8 19 32 2.8 18 19 161 8 20 64 54.5 2.4 17 8.5 23.5 2.8 18 19 161 8 20 64 54.5 3.5 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 725 15.4						
PBPH443 89 954 10.7 PAO 115 1100 9.5 10 1 86 1001 11.6 2 25.5 72 2.8 3 29.5 495 16.8 4 22 70.5 3.2 15 6 20 106 5.3 7 20 37.5 1.9 8 15 31.5 2.1 9 75 1465 19.5 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 16 10 24.5 2.4 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 30 21 54 802 14.9 22 21.5 137.5 6.4 1 24 747 2925 3.9 25 47 725 15.4 26 16.5 61 3.7						
115 1100 9.5 10	•		pHPH443	89		
10			p Δ0	115		
2 25.5 72 2.8 3 29.5 495 16.8 4 22 70.5 3.2 5 39 111.5 2.9 6 20 106 5.3 7 20 37.5 1.9 8 15 31.5 2.1 9 75 1465 19.5 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 16 10 24.5 2.8 16 10 24.5 2.8 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4		10	1	86		
3 29.5 495 16.8 4 22 70.5 3.2 5 39 111.5 2.9 6 20 106 5.3 7 20 37.5 1.9 8 15 31.5 2.1 9 75 1465 19.5 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 16 10 24.5 2.4 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 30 21 54 802 14.9 22 21.5 137.5 6.4 1. 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 27 11.5 16.5 61 3.7			2	25.5		
4 22 70.5 3.2 5 39 111.5 2.9 6 20 106 5.3 7 20 37.5 1.9 8 15 31.5 2.1 9 75 1465 19.5 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 18 32 1.8 15 18 32 1.8 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 30 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4			3	29.5		
5 39 111.5 2.9 6 20 106 5.3 7 20 37.5 1.9 8 15 31.5 2.1 9 75 1465 19.5 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 18 32 1.8 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 30 21 54 802 14.9 22 21.5 137.5 6.4 1.2 23 107 1417 13.2 24 747 2925 3.9 25 46 16.5 61 3.7			• 4	22		
15 6 20 106 5.3 7 20 37.5 1.9 8 15 31.5 2.1 9 75 1465 19.5 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 16 10 24.5 2.4 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 21 54 802 14.9 22 21.5 137.5 6.4 24 747 2925 3.9 25 47 725 15.4 27 18 18 3.9 28 19 16.1 3.7			5	39		
7 20 37.5 1.9 8 15 31.5 2.1 9 75 1465 19.5 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 18 32 1.8 15 18 32 1.8 17 8.5 23.5 2.4 18 19 161 8 19 32 223 7 30 21 54 802 14.9 22 21.5 137.5 6.4 24 747 2925 3.9 25 47 725 15.4 26 16.5 61 3.7		15	6	20		
8 1.5 31.5 2.1 9 75 1465 19.5 20 11 26 95.5 4.75 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 18 32 1.8 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 30 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 27 18.5 12.1 27 18.5 12.1 28.7 14.9 15.1 29 15.4 15.5 15.4 20 15.5 16.5 16.5 17.5 15.4				20		
9 75 1465 19.5 10 20 95 4.75 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 16 10 24.5 2.4 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 21 54 802 14.9 22 21.5 137.5 6.4 1 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 27 18 16.5 61 3.7				15		
20 95 4.75 20 11 26 133.5 5.1 12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 18 32 1.8 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 27 13.5 6.1 28 15.5 61 3.7				. 75		
25			10	20	95	
12 127 1467 11.6 13 33 301.5 9.1 14 317 2280 7.2 15 18 32 1.8 16 10 24.5 2.4 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 54.5 3.5 8.5 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 36 16.5 61 3.7		20	11	26	133.5	
13 33 301.5 9.1 14 317 2280 7.2 15 18 32 1.8 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 27 13.5 61.5 61 3.7			12	127		
14 317 2280 7.2 15 18 32 1.8 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 27 27 325			13	33		
15 18 32 1.8 25 16 10 24.5 2.4 17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 27 27 27			14	317		
16 10 24.5 2.4 17 8.5 23.5 2.8 18 19 161 8 19 161 8 19 32 223 7 20 64 543.5 8.5 22 21.5 137.5 6.4 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 27 27 27 27 27 27 27 27 27 27 27 27 27				18		
17 8.5 23.5 2.8 18 19 161 8 19 32 223 7 20 64 543.5 8.5 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 35 26 16.5 61 3.7		25	16	10	24.5	
18 19 161 8 19 32 223 7 20 64 543.5 8.5 30 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 35 26 16.5 61 3.7				8.5		
19 32 223 7 20 64 543.5 8.5 30 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 35 26 16.5 61 3.7				19		
20 64 543.5 8.5 30 21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 35 26 16.5 61 3.7				32		
21 54 802 14.9 22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 35 26 16.5 61 3.7				64	543.5	
22 21.5 137.5 6.4 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 35 26 16.5 61 3.7		30		54	802	
. 23 107 1417 13.2 24 747 2925 3.9 25 47 725 15.4 35 26 16.5 61 3.7				21.5	137.5	
24 747 2925 3.9 25 47 725 15.4 35 26 16.5 61 3.7				107	1417	
. 25 47 725 15.4 16.5 61 3.7	٠			747	2925	
35 26 16.5 61 3.7 27 33.5	:			47	725	
27	•	35	26	16.5	61	
3.3			27	11.5		

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			180		
	28	13.5	14	1	
	29	44	491	11.2	
5	30	19.5	103.5	5.3	
	31	41.5	182	4.4	•
	32	27.5	125	4.5	•
	33	98.5	1249.5	12.7	•
	34	214.5	2256	10.5	
10	35	69.5	774	11.1	
	36	94.5	1019.5	10.8	
	37	120.5	1073	8.9	
	38	117.5	1790	15.2	
	39	96.5	925	9.6	
15	40	112.5	1501.5	13.3	
	41	97	1085	11.2	
	42	43.5	602.5	13.9	
	43	27.5	298	10.8	
	44	41.5	298	7.2	
20	45	80	994.5	12.4	
	46	42.5	484.5	11.4	
	47	36	328.5	9.1	
	48	40.5	379.5	9.4	
	49	26	166	6.4	
25	50	44.5	349.5	7.9	
	51	57.5	483.5	8.4	
	52	46	328	7.1	
	53	49	469.5	9.6	
	54	52	518	10	
30	55	58.5	471.5	8.1	
	56	31	239.5	7.7	
	57	29	245	B.4	•
	58	32	288.5	9.0	•
	59	38.5	209.5	5.4	•
35	60	31.5	199	6.3	•
	62	23	178	7.7	
			· -	1.1	

		181						
		63	29	268	9.2			
		64	35	196.5	5.6	_		
	5	65	20.5	219.5	10.7			
•		66	26	291.5	11.2			
		67	156.5	1256.5	8.0			
•		68	132.5	1045	7.9			
		69	138.5	1101.5	7.9			
	10	70	397.5	1726	4.3			

 \star GUS activity expressed as flourescence units/hr/10 6 protoplasts.

The results show that, with the exception of the pA28 construction, all 70 modification of the 77 bp element from the 2-2 promoter are able to impart chemical inducibilty to heterologous promoters. It is not known why the pA28 construction was not able

to respond to treatment with N-(aminocarbony1)-220 chlorobenzene sulfonamide

EXAMPLE 21

The Use of N-(aminocarbonyl)-2-chlorobenzene-

30

25 sulfonamide to Induce Expression of the Petunia gene P6.1 in Transgenic Tobacco

The 5' and 3' end mapping data in Example 5 showed that the P614 construction contained a 1.3 kb promoter fragment and a 2.2 kb downstream fragment of the petunia P6 gene. The P614 construction was transformed into tobacco to determine both if this petunia DNA fragment included all the elements necessary for chemical induction, and if this petunia

gene could be both expressed and chemically induced 35 in a heterologous plant species. Plasmid P614 was linearized with Bam HI site and ligated into the Bam

	HI site of the binary vector pAGS135. The binary
	vector pAGS135 used in this example is but one of a
5	large number of binary vectors available that could
	be used for this purpose. pAGS135 is a cosmid binary
	vector whose replicon is derived from the broad host
	range plasmid pRK2 and contains left and right
	borders fragment from the octopine Ti plasmids pTiA6
10	and pTiAch5, respectively [van den Elzen et al.,
	Plant Mol. Biol., 5: 149-154 (1985)]. The border
	fragments delimit the asset (1985)]. The border
	fragments delimit the segment of DNA which becomes incorporated into the host plant genome during the
	process of Agrobatta
15	process of Agrobacterium-mediated transformation. A
	chimeric marker gene (consisting of a neomycinphospho-
	transferase (NPTII) coding region linked to the
	nopaline synthase promoter and the octopine synthase
	3' end) which specifies kanamycin resistance in plant
20	cells is positioned between the left and right border
	fragments. A unique Bam HI site downstream of the
	NPTII gene served as a convenient cloning site. The
	plasmid pAGS135 differs from the plasmid pAGS112
	[disclosed in van den Elzen et al., Plant Mol. Bio.,
25	5: 149-154 (1985)] in that the Xho I in pAGS112
	downstream from the right border has been deleted by
	digestion of pagsil2 with Xho I and ro discuss the
	the plasmid by self-ligation after blunting the Xho I
	overnangs. An aliquot of the lighting.
30	abed to transformed E. coli HB101, and tennes.
	were grown on LB containing ampicillin (75 ug/-1)
	scale placmid
	preparations were made from antibiotic residence
	colonies and digested with to completion with any
25	to identify the colonies with the desired
35	construction. The orientation of the plasmid P614 in
	the binary v ctor (determined by Hind III digests)

15

30

was such that transcription would proceed towards the right T-DNA border, with pucl18 sequences between the end of the petunia gene and the right T-DNA border. This plasmid DNA construction was called P627 (Figure 25).

Transformation of Tobacco with the Petunia P6.1 Gene

The plasmid P627 was moved into Aurobacterium tumefaciens (AL4404/pAL4404) by a triparental mating. Agrobacteria were grown to stationary phase in minimal A medium, while P627 and pRK 2013 (necessary for mobilization of plasmid) were grown for a few hours to logarithmic growth in LB broth. Equal volumes (0.5 ml) of the three strains were concentrated and plated on one LB plate and allowed to grow overnight at 28°C. A loopfull of cells was

- Scraped off the plate and resuspended in 3 ml of 10 mM MgSO₄. Serial ten fold dilutions of these cells (in 10 mM MgSO₄) were plated on LB containing rifampicin (100 µg/ml) and tetracycline (1 µg/ml) and incubated for 3 days at 28°C. Antibiotic resistant colonies were streaked onto minimal A plates
- 25 containing 1 µg/ml tetracycline and incubated for 3 days at 28°c.

Tobacco (SR1) was used as the recipient for transformation. In vitro grown leaf material was sliced into strips using a scalpel. The strips were dipped into <u>Agrobacterium tummefaciens</u> containing the construct P627 (bacterial concentration was 0.2 A550). Leaf pieces were placed on media containing MS major salts, MS minor salts, B5 vitamins, MS iron.

3% sucrose, 0.1 µg/l NAA, 1.0% BA, 0.7% TC agar, pH
35 5.8 and incubated for 2-3 days under growlights.
Leaf material was removed and washed by placing in

liquid culture medium containing 500 µg/1 cefotaxime and rotating gently for 3-4 hours. The leaf pieces were then placed on medium containing 100 µg/1 carbinicillin and 300 mg/l kanamycin and transferred every 2 weeks. Shoots appeared after 2-8 weeks and were transferred to rooting medium (0.5% MS major salts, MS minor salts, iron, 1% sucrose, 0.8% agar, 10 and 2 μM indolebutyric acid). Eight independent transformed plants were regenerated. Plants were transferred to the greenhouse and grown hydroponically in the apparatus described in Example 4 when they became 2-3 inches tall. Two plants that 15 had been regenerated from cell culture, but not transformed were also transferred and included as controls.

Expression of the P6.1 Gene in Transgenic Tobacco 20 Three weeks following the transfer of transformed and control plants to hydroponics, half the exposed roots extending through the foam plug from each of the plants were harvested and frozen in liquid N2. The plants were then treated 25 hydroponically with 200 mg/l N-(aminocarbonyl)-2-chlorobenzenesulfonamide as described in Example 4. After six hours of chemical treatment, the remainder of the exposed roots were harvested and frozen as above. The plants, in their foam plugs, 30 were transferred to soil in shaded pots in the greenhouse for 2-3 days to let roots still in the foam to grow out. Plants were then transferred to the light and grown to maturity. RNA was prepared from root tissue as described earlier. RNAse protection analysis was then performed as 35 described in Example 4 to determine the inducibility

	of the transforming P6 petunia gene as well as the
5	used for this analysis was prepared by digesting the plasmid P611 to completion with Pvu II confidence
10	synthesizing an RNA probe complementary to the coding strand of the P6.1 mRNA using T3 RNA polymerase. The Pvu II site occurs 150 bp from the 3' end of of the Eco RI fragment in P611 and therefore should genreate a protected fragment of 150 bp if the introduced
15	petunia gene is expressed in tobacco. All eight transformants demonstrated inducible expression of the transferred gene in their roots (Figure 26). These results demonstrated that the 4.5 kbp petunia genomic DNA fragment contained all the elements required for induction of the gene by N-(aminocarbonyl)-2-chlorobenzenesulfonamide, and that this
	inducibility could be transferred to another species

20

Expression of the P6.1 Gene in Transgenic Tobacco Callus

The inducibility of the P6.1 gene was also examined in callus tissue derived from transformed 25 tobacco plants. It was felt that if the expression of chemically inducible genes were responsive to chemical stimulation in callus, then testing and selection for callus to be regenerated to whole plants could be accelerated. To this end, leaf 30 tissue from one of the the P6.1 tobacco transformants was placed on media that supports callus induction (MS media containing 0.1 $\mu g/l$ napthalene acetic acid and 0.3 µg/l kinetin). After five weeks, 1-1.5 cm calli had developed. These calli were transferred to liquid media (MS media containing 0.1 μ g/1 napthalene acetic acid and 0.1 µg/l benzyladenine) and shaken at

15

20

25

28°C overnight. The next day, pieces of the callus
were transferred to MS media, or MS media containing
100 mg/l N-(aminocarbonyl)-2-chlorobenzenesulfonamid
and shaken at 28°C. Samples of callus tissue were
removed from the flasks at 6 and 20 hours and frozen
in liquid N. DNN
in liquid N2. RNA was prepared from callus tissue
using the procedure described in Example 4. The
inducibility of both the introduced petunia P6 gene
and the endogenous tobacco T2.1 gene by N-(amino-
carbonyl)-2-chlorobenzenesulfonamide treatment was
evaluated using the RNAse protection analysis as
described above. Both the endogenous tobacco gene
and the transforming petunia gene were barely
detectable in untreated callus tissue, while strong
expression of both genes was observed in
N-(aminocarbony1)-2-chlorobenzenesulfonamide-treated
callus. The levels of expression observed for both
genes approximated that seen for their expression in
the root tissue of intact, chemically treated tobacco
plants. It was therefore concluded that the
inducibility of foreign genes whose expression is
regulated by promoters responsive to substituted
benzenesulfonamides is assayable at the level of
assayable at the level of

EXAMPLE 22

30 The Use of N-(aminocarbonyl)-2-chlorobenzenesulfonamide to Induce Expression of Recombinant Petunia gene P6.1 promoter/GUS fusions in Trangenic Tobacco Plants

35 Construction of P655

transformed callus tissue.

The plasmid P655 was digested to completion with Hind III and Bam HI and the resulting DNA

fragments were separated by agarose gel electrophoresis. The 3.9 kbp DNA fragment.

- 5 containing a recombinant gene consisting of a GUS coding region operably linked to a 1.3 kbp P6.1 promoter fragment and an OCS 3' downstream region, was excised from the gel and recovered by electroelution as described earlier. The DNA was
- then extracted with an equal volume of phenol:chloroform (1:1 v/v) and ethanol precipitated. The binary vector pAGS502 was digested to completion with Hind III and Bam HI, extracted
- with an equal volume of phenol:chloroform (1:1 v/v)

 and ethanol precipitated. Equimolar amounts of
 vector and the gel purified 2.0 bb. de-
- vector and the gel purified 3.9 kbp insert were ligated in 10 µl for 4 hours at 15°C. An aliquot of the ligation mixture was used to transform E. coli HB101 and aliquots of the resulting transformed cells
 - were plated on LB plates containing 10 µg/ml tetracycline. Small scale plasmid preparations were prepared from tetracycline-resistant colonies and subjected to digestion with Hind III and Bam HI until a colony was found that contained the desired 3.9 kbp
 - DNA fragment in the binary vector pAGS 502. The binary vector pAGS502 used in this example is but one of a large number of binary vectors are available and could be used for this purpose. To make pAGS502 the Eco RI-Hind fragment of pAGS111 [van den Elzen et
 - 30 al., Plant Mol. Biol., 5: 149-154 (1985)] (consisting of a NOS/NPTII/OCS 3' end gene between the left and right T-DNA borders) was rendered blunt and cloned
- into the blunted Eco RI site of the wide host range plasmid pRK290 [disclosed in Ditta et al., Proc. 35 Natl. Acad. Sci. USA., 77: 7347-7351 (1980)]. The
 - Xho I site downstream from the right border was

15

deleted by digestion with Xho'I and re-circularizing
the plasmid by self-ligation after blunting the Xho I
5' overhangs. The polylinker sequence 5'GGATCCTCTAGAAAGCTTCGAACTCGAGGAATTCGTT-3' was then
inserted between the Bam HI-Hpa I sites within the
T-DNA borders to create pAGS502. This plasmid

construction was designated P656 (Figure 27).

Construction of P661

The methods used to create the plasmid construction P656 were repeated using the plasmid constructions P658 and pAGS 502 as starting materials: A 4.7 kbp DNA fragment, consisting of a GUS structural gene operably linked to a 600 bp P6.1 promoter fragment and a 2.2 kbp P6.1 3 end fragment, results from digestion of P660 with Hind III and Bam

HI. This 4.7 kbp DNA fragment was subcloned into Bam HI/Hind III digested pAGS502 as described above and the resulting plasmid construction was designated P661 (Figure 27).

Construction of P662

25 The methods used to create the plasmid construction P656 were repeated using the plasmid construction P658 were repeated using the plasmid construction P658 as the starting material. A 4.4 kbp DNA fragment, consisting of a GUS structural gene operably linked to a 300 bp P6.1 promoter fragment and a 2.2 kbp P6.1 3' end fragment, results from digestion of p658 with Hind III and Bam HI. This 4.4 kbp DNA fragment was subcloned into Bam HI/Hind III digested pAGS502 as described above and the resulting plasmid construction was designated P662 (Figure 27).

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Construction of P663

The methods used to create the plasmid Construction P656 were repeated using the plasmid construction P657 and pAGS 502 as the starting materials. A 5.4 kbp DNA fragment, consisting of a GUS structural gene operably linked to a 1.3 kbp P6.1 promoter fragment and a 2.2 kbp P6.1 3' end fragment, results from digestion of P658 with Hind III and Bam HI. This 5.4 kbp DNA fragment was subcloned into Bam HI/Hind III digested pAGS 502 as described above and the resulting plasmid construction was designated P663 (Figure 27)

Transformation of Tobacco with P661, P662, and P663

The plasmids P656, P662, and P663 were moved into <u>Agrobacterium tumefasciens</u> (AL4404/pAL4404) using the triparental mating procedure and tobacco

20 (SR1) leaf pieces were transformed with each of the four chimeric GUS/P6.1 fusions using the procedures described in Example 21.

Induction of GUS Activity by N-(aminocarbonyl)-2-chlorobenzenesulfonamide

A number of regenerated plants that had been transformed with the P661, P662, or P663 constructions were transferred to the hydroponic system described in Example 5. Root tissue was harvested from these hydroponically-grown plants and treated with N-(aminocarbony1)-2-chlorobenzenesulfonamide as described in Example 14. The root material was then used to make crude protein extracts which were tested for GUS activity. The plants were

35 then transferred to soil in pots and grown to maturity in a greenhouse as described earlier.

		Roots were homogenized in ice cold GUS assay	
		Duffer (50 mM sodium phosphate pH 7.0, 10 mM pro	
•	5	U.1% Triton X-100, 1mM EDTA using a Dounce type	
		nomogenizer. Cellular debris was then removed by	
		centrifugation. Fluorometric GUS assays were	
		performed using a Perkin-Elmer Fluorescence	
		Spectrophotometer (650-40) set for an excitation	
	10	wavelength of 365 nm and an emission wavelength as	
		455 nm. A standard fluorescense vs. MU concentration	
		curve was prepared by diluting 50 ml of various	
		concentrations of MU into 950 µl 0.2 M Na ₂ CO ₃ and	
		measuring the fluorescence.	
	15	GUS activity in root extracts of transformed	
		plants was assayed by adding 15 µl of the substrate	
		(1 mM 4-methyl umbilliferyl glucuronide in assay	
		buffer) to 1 ml of crude root extract and incubating	
		at 37°C. Fluorescence measurements were taken at of	
	20	0, 15 and 30 minute time points by adding a measured	
		amount (1 to 50 µl) of the GUS reaction to 1 ml 0.2 M	
		Na ₂ CO ₃ and measuring the fluorescence of the MU	
		generated in the GUS reaction. Protein	
		concentrations in the crude root extracts were	
	25	determined by Bradford protein assays. From 10 to 20	
		μl of root extract were added to 1 ml of Bradford	
		Assay Stain (10 µg/ml Coomassie Brilliant Blue G in	
		8.5% phosphoric acid) and the absorbance of the	
		samples was measured at 595 nm. A protein	
	30	concentration vs. absorbance curve was prepared using	
		BSA as a protein standard. GUS activity in each root	
		extract was standardized to protein concentration and	
		expressed as GUS activity per microgram protein.	:
		The results of on such analysis are shown in	_
	35	Table 10. A number of plants transformed with the	·
		P661, P662, and P663 constructions show induction of	
		and a show induction of	

GUS activity following treatment with N-(amino-Carbonyl)-2-chlorobenzenesulfonamide. The

5 variability seen in the expression of the chimeric GUS gene is commonly seen when testing primary transformants for the expression of a transforming

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TABLE_10

N-(aminocarbony1)-2-chlorobenzenesulfonamide Inducible GUS Expression in Tobacco Plants Transformed with Petunia P6 Promoter/Gus Constructions

15

Plant #	Promoter Size (in bp)	GUS ACT (FU/µg- Uninduced	IVITY min) Induced	Fold Induction
P663/10 P663/11 P663/17 P663/36 P663/81 P661/105 P662/44 P662/55 P662/65	1300 1300 1300 1300 600 600 300 300	7.4 3.1 6.7 8.5 31.5 45.8 4.3 2.2 25.8 2.9	13.6 8.9 21.0 44.9 31.2 102.1 7.0 7.8 76.6 5.1	1.8 2.9 3.1 5.3 1.0 2.2 1.6 3.5 3.5

EXAMPLE 23

The Use of N-(aminocarbonyl)-2-chlorobenzenesulfonamide to Induce Expression of Recombinant Corn 2-1 Promoter/GUS Gene Constructions in Trangenic Tobacco

Construction of pJE518 and pJE519

The recombinant 2-1/GUS gene contained in plasmid pJE516 was stabily introduced into tobacco by Agrobacterium mediated transformation. The plasmid pJE516 was digested to completion with Bam HI and Xho I

35 and the resulting 6.0 kbp DNA fragment consisting of a $3\,$

	kbp 2-1 promoter fragment/GUS/1.1 kbp of 2-1 gene 3'	
5	downstream fragment was gel purified mb/-	
3	putitied 6 kb Bam HI/Xho I fragment from - The con-	
	and rigated into the Bam HI/Yho I ait	:
-	The binary vector pilaces	
	and a saige number of binary vectors that	:
	available and may be used in this example. It was	
10	derived from the broad host range vector pRK2 and	
	contains a hygromycin resistance gene (HYG) under	
	control of the Agrobacterium 1',2' promoter and	
	nopaline synthese 21 and to	
	nopaline synthase 3' end between the left and right	
15	T-DNA borders. The HYG gene specifies hygromycin	
	resistance in transformed plants. A polylinker	
	sequence was inserted downstream from the HYG gene to	
	rioride a set or unique restriction site for	
	cloning. The Xho I site downstream of the T-DNA	
20	right border was removed as described	
	phobboz. The resulting plasmid was dealers	
	r (rigule 28).	
	The plasmid pJE 516 was also digested to	
	completion with Bam HI and Hoa I This	
	The liagment from the vector gonei-ti-	
25	Promoter fragment/GUS/1 1 kbp as a	
	Transcreding Iragment fusion. This factor	
	Further and ligated into the Ram DIAU	
	to create the plasmid nJFS10 /Pi	
	These two plasmid construction were used to transform	
30	tobacco (Petite Havanna).	

Transformation of Tobacco

35

The constructs in pJE518 and pJE519 were mobilized from E. coli HB101 into Agrobacterium tumefaciens in order to perform tobacco transformation. Fresh cultures of Agrobacterium

			AL4404 harboring plasmid pAL4404 were grown in
_			Minimal A media (10.5g K ₂ PO ₄ , 4.5g KH ₂ PO ₄ , ug
	:	5	(NH ₄) ₂ SO ₄ , 0.5 g NaCitrate · 2H ₂ O, 1 ml 1M MgSO. · 7H O
			10 ml 20% glucose, water to 1 1). E. coli HB101
	:		harboring plasmid pRK 2013, and E. coli HB101 strains
			harboring the plasmids to be mobilized (pJE518 and
			DJE519) were grown even-interest
		10	numbers of each type of cells were mixed together,
			plated on LB plates, and allowed to grow at 28°C
			Overnight A loop full of the
			overnight. A loop full of the resulting bacteria was
			suspended in 10 mM MgSO ₄ , plated at 10^0 , 10^{-2} and 10^{-4} dilutions of 10^{-4
		15	10-4 dilutions on LB plates with 100 µg/ml
			rifampicin, 1 µg/ml tetracycline and allowed to grow
			at 28°C for 2-3 days. Single colonies growing on
			these plates were streaked on minimal A plates
			(minimal A media plus 1% agar) containing 1 µg/ml
		20	tetracycline. Overnight liquid cultures were grown
		20	from these streaked colonies in minimal A at 28°C.
			Leaves were taken from 3-4 inch tall tobacco
			plants (Petit Havana) that had been grown in Magenta
			boxes and cut crosswise into approximately 5 mm wide
			strips using a surgical scalpel. The strips were
		25	then dipped briefly into the agrobacterium overnight
			Culture and placed on bacterial cocultivation
			plates. Bacterial cocultivation plates contain Mc
			salts (1.9 g/1 KNO3, 1.65 g/1 NH4NO3, 0.44 g/1
			CaCl ₂ ·2H ₂ O, 0.37 g/l MgSO ₄ ·7H ₂ O, 0.17 g/l KH ₂ PO.
		30	10.3 mg/1 ZnSO ₄ · 7H ₂ O, 16.9 mg/lMnSO ₄ · H ₂ O, 6.2 mg/l
			H ₃ BO ₃ , 0.84 mg/l KI, 0.2 5mg/l Na ₂ MoO ₄ • 2H ₂ O ₄ , 0.025
	÷		pg/1 CusO ₄ · 5H ₂ O, 0.025 mg/1 CoCl ₂ · 6H ₂ O, 37.2 ug/1
			Na ₂ EDTA·2H ₂ O, 27.8 μg/l FeSO ₄ ·7H ₂ O), B5 vitamins (1
	:		μg/l nicotinic acid, 10 μg/l thiamine HCl, 1 μg/l
		35	pyridoxine HCl, 100 µg/l myo-inositol), 0.59 g/l MFC

	227	
	30 g/l sucrose, 8 g/l agar, 0.1 µg/l napthaleneacetic	
5	dela, and i μg/I benzyladenine.	*
	(same media as used in bacterial co-cultivation	:
10	plates, but without agar) containing 500 µg/l cefotaxime. The leaf pieces were then placed on MS medium containing 100 µg/l vancomycin and 30 µg/l	
15	any tomycin and incubated at 27°C and incubated under the same conditions described above. Shoots began to appear after about one month. These shoots were transferred to MS media containing 1 mM indolebutyric acid and 30 mg/l hygromycin when they were about 1 cm tall. Plantlets were moved to Magenta boxes (containing the same media), and allowed to make	
20	2-3 inches tall before being moved to hydroponics.	
25	Induction of GUS Activity by N-(aminocarbony1)-2- chlorobenzenesulfonamide Seven plants transformed with the pJE518 construction and five plants transformed with the pJE519 construction were transferred to the hydroponic system described in Paramile	

construction and five plants transformed with the

25 pJE519 construction were transferred to the
hydroponic system described in Example 4. These
transformants were grown hydroponically until they
had developed sufficient root mass to allow removal
of small samples without destroying the plants. At
this point approximately one-third of the root
material from each plant was harvested and frozen in

material from each plant was harvested and frozen in liquid nitrogen. The plants were then moved to trays containing 0.5% Hoagland's solution with 200 mg/1 N-(aminocarbony1)-2-chlorobenzenesulfonamide. After

35 6 hours of incubation in the presence of the chemical, another one-third of the root material was

harvested from each plant. Root material was used to make crude protein extracts which were tested for GUS

activity. Plants were then transferred to soil in pots and grown to maturity in a greenhouse.

Root material was homogenized in ice cold GUS assay buffer (50 mM sodium phosphate pH 7.0, 10 mM DTT, 0.1 % Triton X-100, 1 mM EDTA using a Polytron (Brinkmann Instruments) GUS activity in roots was then measured after 0, 1, 2, and 4 hours as described in Example 15.

The results of this analysis are shown in Table
11. A number of plants transformed with the pJE 518
15 and the pJE519 construction show up to a 13 fold
induction of GUS activity following treatment with
N-(aminocarbony1)-2-chlorobenzenesulfonamide. The
variability seen in the expression of the recombinant
2-1 promoter/GUS construction is commonly seen when

- 20 testing primary transformants for the expression of a transforming gene. The plants showing the highest level of responsiveness to chemical treatment were both self-fertilized and backcrossed to Petite Havana tobacco. Seeds resulting from backcrosses of a
- number of these plants were germinated and grown in Magenta boxes with a hygromycin selection. After a root structure formed on each plant, root pieces of each were excised and incubated overnight on rooting media with or without N-(aminocarbony1)-2-chloro-
- benzenesulfonamide. GUS assays were performed on extracts of these roots on the following day. The results of this assay are given in Table 12. Roots from the progeny of the backcrosses show N-(aminocarbony1)-2-chlorobenzenesulfonamide inducible GUS
- 35 activity, with two plants transformed with the pJE519 construction showing a ten-fold induction.

		Additional progeny of these crosses will be tested for the inducibility of the recombinant GUS gene in	
	5	response to the both hydroponic and foliar	
		application of N-(aminocarbony1)-2-chlorobenzene- sulfonamide.	
		:	
	10		
	15		
:	20		

BLE	

N-(aminocarbonyl)-2-chlorobenzenesulfonamide Inducible GUS Expression in Tobacco Plants Transformed with 2-1 Promoter/Gus Constructions

A. Transgenic plants containing pJE518 construction

GUS ACTIVITY (FU/ug-min)

	Plant #	Uninduced*	Induced*	Fold Induction
15	518-1 518-2 518-3 518-4 518-5 518-6 518-7	1.0 1.2 0.67 2.9 0.72 0.74	1.9 1.8 8.9 5.5 0.24 2.4 5.0	1.9 1.5 13.0 1.9 0.33 3.2 7.5

B. Transgenic plants containing pJE519 construction GUS ACTIVITY (FU/µg-min)

25	Plant # 519-1 519-2 519-3 519-4 519-5	Uninduced* 0.75 0.70 0.41 1.56	1.8 1.1 1.3 7.2	Fold Induction 2.4 1.5 3.2
	519-5	1.56 0.39	7.2 3.9	4.6 10.0

*Induction in Table 7 was accomplished by hydroponic treatment transformed plants with 200 mg/l of 30 N-(aminocarbony1)-2-chlorobenzenesulfonamide

5

TABLE 12

GUS Assays for Backcross Progeny
GUS Activity
(FU/mg-min)

Fold 10 Fold Induction Plant Cross -D5293 +D5293 Induction of Parent 801-3 518.2 X 1.04 4.4 4.2 1.5 Petite Havana 801-4 1.03 5.3 5.1 1.5 15 802-5 5.8.6 X 1.39 7.9 5.7 3.2 Petite Havana 802-6 1.46 4.8 3.3 3.2 803-5 519.3 X 0.20 2.0 10 3.2 Petite Havana 20 803-6 0.16 1.5 9.4 3.2

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EXAMPLE 24

The Use of N-(aminocarbonyl)-2-chlore-

5 benzenesulfonamide to Induce Expression of Recombinant Genes Under the Control of the Corn 2-2 Promoter in Transgenic Tobacco

Construction of pJE573, pJE578-1 and pJE578-8

The chimeric 2-2/GUS gene contained in plasmid pTDS130 was stabily introduced into tobacco by agrobacterium mediated transformation. The plasmid pTDS130 was digested to completion with Xba I and the

- 15 resulting 4.3 kbp DNA fragment consisting of a 1.2 kbp 2-2 promoter fragment/GUS/1.1 kbp of 2-1 gene 3 downstream fragment was gel purified. This purified 4.5 kbp Xba I fragment from pJE516 was then ligated
- into the binary vector pages502 to yield the plasmid
 pJE573.
 The plasmid pTDS130 was also digested to

completion with Bam HI and the 3.4 kbp DNA fragment consisting of a 0.45 kbp 2-1 promoter fragment/GUS/1.1 kbp of 2-1 gene 3' downstream fragment from pJE516 was then ligated into the binary

vector pAGS502 to yield the plasmids pJE578-1 and pJE578-8. These two plasmids represent each of the two possible orientations of the 2-2 recombinant constructions in the binary vector.

Construction of pDuPU3

The chimeric 2-2/HRA gene contained in plasmid pDuFS22 was digested with Kpn I and Sal I and the resulting 4.3 kbp DNA fragment consisting of a 0.45 kbp 2-2 promoter fragment/HRA/1.1 kbp of 2-1 gene 3

downstream						This	purified
Eragment fr							
the binary	vector p	2596	+0 1	11011	6ha -1		

Transformation of Plants

Mobilization of the constructs in pJE573,

10 pJE578-1 and pJE578-8 from E. coli (strain HB101)

into Agrobacterium tumefaciens, transformation of SR1
tobacco leaf disks, and regeneration of plants were
performed as described in Example 23.

The recombinant ALS gene in pDuPU3 was

15 mobilized from E. coli HB101 into Agrobacterium
tumefaciens strain LBA4404 by a triparental mating.
E. coli HB101 containing the plasmid pRK2013 was used
as a helper for plasmid mobilization matings.

Bacterial strains HB101 containing pDUPU3. HB101 with

pRK2013, and LBA4404 were grown overnight in 5 ml of LB broth with appropriate selective antibiotics.

Bacteria were harvested by centrifugation at 4000X g for 10 minutes at 22°C and resuspended in 5 ml LB broth. Metings were performed by mixing 100 ul of

each culture in a 1.5 ml microfuge tube and pipetting aliquots of the mixture onto sterile Millipore type HA nitrocellulose disks. Disks were placed on 6-8 sheets of sterile Whatman #1 filter paper to remove excess liquid from cultures and then transferred to

30 LB agar in 100 mm petri dishes. After incubation for approximately 16 hours at 30°C, bacteria were washed from the nitrocellulose discs into sterile 4 ml polypropylene culture tube using 1 ml of 10 mM MgSO₄. The bact ria were serially diluted and

35 various dilutions were plated onto LB agar plates containing 100 ug/ml each of rifampacin and ampicillin and incubated at 30°C. Small scale

plasmid preparations made from resistant colonies

were analyzed for the presence of the desired income

5 were analyzed for the presence of the desired insert DNA by Southern blot analysis of Ti plasmids.

Transformation of Plants

- Constructions were introduced into the plants
 via Agrobacterium tummifaciens infection of tobacco
 leaf disks. Standard aseptic techniques for the
 manipulation of sterile media and axenic
- plant/bacterial cultures were followed, including the
 15 use of a laminar flow hood for all transfers. Potted
 tobacco plants for leaf disk infections were grown in
 a growth chamber maintained for a 12 hour, 24°C day,
 12 hour, 20°C night cycle, at 80% relative humidity,
 under mixed cool white fluorescent and incandescent
- 20 lights. Tobacco leaf disk infections were carried out essentially by the method of Horsch et al. (Horsch, R. B., Fry, J. E., Hoffman, N. L., Eichholtz, D., Rogers, S. G., and Fraley, R. T. (1985) Science 227, 1229-1231).
- 25 Young 4-6 inch partially expanded leaves were harvested with a scalpel from 4-6 week old plants. The leaves were surface sterilized for 30 minutes by submerging them in approximately 500 ml of a 10% Clorox, 0.1% SDS solution and then rinsing 3 times
- 30 with sterile deionized water. Leaf disks were then prepared using a sterile paper 6 mm punch and they were inoculated by submerging them for several minutes in 20 ml of a 1:10 dilution of an overnight LB broth culture of Agrobacteria carrying the plasmid
 - of interest. After inoculation, leaf disks were placed in petri dishes containing CN agar medium (MS

salts	(Gibco)	30	gm	sucrose,	8	gm	agar,	0.1	m1	of	1
mg/ml	NAA, and	11	ml	of 1 mg/r	11	BAI	per	liter	, F	H	_

5 5.8). The plates were sealed with parafilm and incubated under mixed fluorescent and "Gro and Sho" plant lights (General Electric) for 2-3 days in a culture room maintained at approximately 25°C.

Leaf disks were transferred to fresh CN medium containing 500 mg/L cefotexime and 100 mg/L kanamycin. The disks were incubated under the growth conditions described above for 3 weeks and then transferred to fresh media of the same composition. Approximately 1-2 weeks later shoots which developed on kanamycin-selected explants were excised with a sterile scalpel and planted in A medium (MS salts (Gibco), 10 gm sucrose, and 8 gm agar per liter) containing 100 mg/L kanamycin. Shoots which rooted

were transferred to soil and grown in a growth
chamber as described above.

Induction of GUS Activity In Plant Transformed with B-glucoronidase Gene Constructions

- 25 Plants transformed with the JE573, pJE578-1 and pJE578-8 constructions were grown hydroponically, treated with 200 mg/l N-(aminocarbonyl)-2chlorobenzene and assayed for the induction of GUS activity as described in Example 23.
- The results of this analysis are shown in Table

 13. A number of plants transformed with the JE573,
 pJE578-1 and pJE578-8 constructions display the
 induction of GUS activity following treatment with
 N-(aminocarbonyl)-2-chlorobenzenesulfonamide. The
- 35 variability seen in the expression of the chimeric GUS gene is commonly seen when testing primary

_	transformants for the expression of a transforming gene. The plants showing the highest level or
;	response to chemical treatment have been selfed, and progeny of these selfs will be tested for stability of the gene and the inducibility of the chimeric GUS gene in response to the foliar application of inducing compounds.

TABLE 13

				LE 13			
5	Plant I.D.	Aci	pecific tivity cotein/min)	Fold Induction	X Progeny Segregatio	#of	:
		UCT 573			Kan-R/Kan-	1 10 <u>01</u> S	•
10	4 5 6	0.54 1.25 0.50	1.12 3.62 3.06	2.1 2.9 6.1	90/11 39/16 102/4	2	·
15	8 9 10 11 12	0.45 0.016 3.73 0.42	1.76 0.128 5.34 1.40	4.0 8.0 1.4 3.3	38/13 0/50 100/18 36/3	3 1 0 [^] 2 2	
		0.35 CT 578-1	0.003	4.4	63/6	2	
20	14 19 27 32	0.003 0.094 0.062 0.272	0.003 0.128 0.127	1 1 1.4 2	41/22 NA 118/4 45/19	1 NA >2 1	
25	35 37 CONSTRUC	0.004 0.018	1.445 0.002 0.145	5.3 1 8.3	53/24 NA 52/16	1 NA 1	
30	6A 6B 7 9	1.86 1.18 0.79 1.45 4.83	6.19 2.73 1.80 4.20 6.25	3.3 2.3 2.3 2.9	46/14 80/35 125/0 NA 121/8	1 1 >2 NA 2	į

Induction of Herbicide-Resistant ALS in Plants Transformed With pDuPS22

Plants transformed with the pDuPS22 construction were grown in soil for three weeks and two upper leaves were harvested from each plant. One leaf was placed into a beaker containing 0.5%

- Hoagland's solution such that the bottom 2 cm of the cut end of the leaf was submerged in liquid. The second leaf was placed in a beaker containing 0.5% Hoagland's containing 200 mg/l N-(aminocarbonyl)-2 chlorobenzenesulfonamide. Leaves were then incubated
- on the growth chamber for 16-24 hours and divided in half. One half was analyzed for the expression of ALS mRNA, while the other was analyzed for the expression of sensitive and herbicide resistant ALS
- The expression of stable cytoplasmic mRNA
 transcribed from the wild type and transforming ALS
 genes in transformed plants were measured by RNAse
 protection analysis. In this manner, expression of
 the pDuPS22 construction was distinguished from the
 wild type ALS genes by virtue of the fact that the

enzyme levels

- ppurs22 transcript has a 2-2 untranslated leader that is divergent from the untranslated leader of the native ALS genes. To this end, the Eco RI/Nco I fragment of the tobacco SurB ALS gene that spans the region from 133 bp 5' to the SurB translation start
 - region from 133 bp 5' to the SurB translation start site to 348 bp beyond the SurB translation start was cloned in to the vector PTS64 to create the plasmid designated pTSNTC (the isolation of the wild type SurB gene is taught in European Patent application
- 35 number 0257993, and a herbicide-resistant SurB gene is available from ATCC as accession number 67124 and

may be	substituted for	the wild-type	SurB gene to
obtain	the same result	.) The plasmi	pTS64 was

- prepared by digesting the plasmid pSP64 (Promga Biotech, Inc.) to completion with Bam HI and ligating the vector with synthetic double stranded oligonucleotide of the sequence
- 5'-GATCTATCGATCCATGGTCTAGAAAA-3' 10
- 3'-ATACGTAGGTACCAGATCTTTT-5'. The ligation mixture was then heated to 65°C for 10 min. and digested to completion with Xba I. The digestion mixture was heated to 65°C for 10 min. again and subjected to ligation with T4 DNA ligase overnight. 15 Following transformation of the ligation mixture into
- competent E. coli DH5, a colony was identified that contained desired sequence
 - 5'-GATCTATCGATCCATGGT-3'
- 3'-ATACGTAGGTACCAGATC-5' encoding a Cla I site and an Nco I site inserted into the pSP64 polylinker. 20

A 520 b 32p-labelled antisense ALS RNA probe was prepared from Eco RI linearized pTSNTC using SP6 polymerase in the presence of α -32p dCTP with a kit by following the manufacturer's

- recommended protocol. Hybridization of wild type ALS 25 mRNA to this 520 b 32P-labelled antisense RNA should protect 410 b of the probe, while hybridization to the pDUPS22 transcript should protect only 348 bp of the probe corresponding to the region 3' to the
- 30 translation start site of the pDuPS22 mRNA. RNAse protection assays were carried out using the protocol of Zinn et al. (Zinn et al. Cell (1983) 34, 865-879). Labelled antisense strand RNA
- was annealed to total FNA from either wild type 35 tobacco plants or to 10 ug of total RNA from plants
 - transformed with the pDuPS22 construction. The sizes

of the labelled RNA fragments remaining after digestion with RNAse Tl and RNAse A were determined by electrophoresis using 6% denaturing polyacrylamide gels. Results of such analyses showed that N-(amino-

	•		gets. Results of such analyses showed that N-(amino-
			Carbonyl)-2 chlorobenzenesulfonamide treatment -
	•		plants transformed with the pDuPS22 2-2 promoter up-
			recombinant gene resulted in the induction of him
		10	levels of stable cytoplasmic HRA mRNA.
			As a preliminary test of the inducibility
			of the sulfonylurea-resistant ALS gene, several small
			leaves were excised from each of sixteen
			kanamycin-resistant shoots, sliced into 2-3 mm
		15	pieces, and placed on callus induction medium that
			consisted of MS salts, 100 mg/L i-inositol, 0.4 mg/L
			thiamine, 3% sucrose, 1 mg/L NAA, 0.2 mg/L BAP, 0.8%
			agar, 500 mg/L cefotaxime, pH 5.8 containing either
_			10 ppb chlorsulfuron, 10 ppb chlorsulfuron+100 ppm
		20	D5293, 100 ppm D5293, or no selective agent. Callus
			formation was scored as plus or minus after three
			weeks of growth. Results are summarized below:
			10/16 formed callus
		25	10 ppb Chlorsulfuron 12/16 formed callus 10 ppb Chlorsulfuron
			100 proce
			0/16 formed callus
			Protein extracts were prepared from leaves of a
		30	number of kanamycin resistant plants that were
			treated with N-(aminocarbonyl)-2 chlorobenzene-
			sulfonamide and assayed for ALS enzyme activity as
	:		described by Chaleff and Mauvais [Chaleff R. C. and
			Mauvais C. J. (1984) Acetolactate synthase is the
	;	35	site of action of two sulfonylurea herbicides in
			higher plants. Science 224:1443-1445]. The reaction
			The reaction

	product, acetoin, was quantified by measuring optical
	density at 530 nm [Westerfield WW (1945) a
	colorometric determination of blood acetoin 7
	Biol. Chem. 161:495-502]. For each extract,
	replicate enzyme assays were performed reactions
	either with no herbicide or 100 pph chloreulaung
	The average ALS activity in the presence of
1	chlorsulfuron, expressed as a percentage of the total
	average ALS activity measured in the absence of
	herbicide, is presented in Table 14.
	These results show that two of the seven plants
	showed increases in the level of chlorsulfuron-
15	resistant ALS following chemical treatment. It
	should be noted that there is a well documented
	biological mechanism that keeps the ALS specific
	activity fixed in tobacco. Therefore, even though
	all plants tested showed induction of herbicide-
20	resistant ALS mRNA, the inability to increase the
	total ALS activity in leaves is to be expected.
	Those plants showing near 100% resistant ALS activity
	when uninduced represent plants where sufficient
	expression of the resistant ALS gene was obtained in
25	the absence of chemical treatment to yield
	significant amounts of resistant enzyme. The level
	of gene expression in untreated plants transformed
	with genes driven by the 2-2 promoter is a position
	effect, and is seen to vary dramatically from
30	undetectable to very high level, both with 2-2/ALS
	and 2-2/GUS gene constructions. It is expected that
	a number of plants with no uninduced ALS activity
	will be found when a larger population of 2-2/ALS
	transformants is studied.
35	

TABLE 14

_			OD 530	OD530 100 ppb	Uninhibited	
		5 Plant	No Herbicide	Chlorsulfuron	Activity	
	•	Untransformed				
		Untreated	0.204	0.010	5	
	•	Treated with D52	93 0.267	0.034	13	
		Transformant #44B				
		10 Untreated	0.333	0.306	92	
		Treated with D529	93 0.385	0.365	95	
		Transformant #53A			•••	
		Untreated	0.244	0.251	103	
		Treated with D529	3 0.331	0.312	94	
		15 Transformant #61A			,,	
		Untreated	0.376	0.347	92	
		Treated with D529	3 0.912	0.901	99	
		Transformant #63A	•		**	
		Untreated	0.457	0.178	39	
		20 Treated with D529	3 0.835	0.732	88	
		Transformant #74C			••	
		Untreated	0.859	0.822	96	
		Treated with D529	3 0.400	0.408	102	
		Transformant #79A				
		25 Untreated	0.492	0.309	63	
		Treated with D529	3 0.366	0.325	89	
		Transformant #93A			0,5	
		Untreated	0.324	0.313	97	
		Treated with D529	0.989	1.003	101	
		30				

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7 N D T T T T E

	EXAMPLE 25
	The Use of N-(aminocarbonyl)-2-chloro-
5	benzenesulfonamide to Induce Expression of a
	Recombinant 2-1 Promoter/GUS Construction in
	Transgenic Brassica
	Standard aseptic techniques for the
10	manipulation of sterile media and axenic
	plant/bacterial cultures were followed, including the
	use of a laminar flow hood for all transfers.
	Seeds of Brassica napus cv. Westar were
	sterilized by soaking in 70% ethanol for three
15	minutes followed by a 20 min treatment in 20% v/v
	bleach (sodium hypochlorite). The seeds were rinsed
	in sterile distilled water three times and planted at
	a density of nine seeds per Magenta box on seed

germination media (Germination media: MS (Murashige 20 and Skoog) salts, 1% sucrose, 3 mM MES buffer, and 0.8% Hazleton TC agar). Seeds were germinated at 24°C using a 16 h light/8 h dark photoperiod with a light intensity of 4000 lux. After five days, the hypocotyls from the germinated seedlings were excised and cut into sections ranging in length from 0.5 to 1.0 cm.

Single colonies from freshly streaked plates of Δ . timefaciens strain LBA4404 containing pJE519 (Example 23) were grown overnight in minimal Λ medium (10.5 g/1 K_JHPO_4, 4.5 g/1 KH_2PO_4, 1.0 g/1 (NH)_2SO_4, 0.5 g/1 Na citrate 2H_2O, to 990 ml; autoclave and add sterile solutions are added; 1 ml of 1 M MgSO_4, 10 ml of 20% glucose. The host strain LBA4404 is rifampicin resistant and the introduced binary plasmid specifies bact rial tetracycline resistance.

The agrobacterium suspensions were diluted in hor<u>mone-free plant media (MS salts, Gamborg's BS</u>

- 5 vitamins, 3% sucrose, 3 mM MES buffer, pH 5.8) to a concentration of 2.8 x 10⁸ cfu/ml using the optical density of the culture at 550 to estimate the bacterial concentration.
- The hypocotyl sections were individually dipped in the agrobacterium suspension and then placed onto sterile Whatman #1 filter paper which had been placed on top of callus regeneration media (MS salts, B5 vitamins, 3% sucrose, 3 mM MES buffer, 0.2 mg/1 2,4-D, 3 mg/1 kinetin, 0.8% Hazleton TC agar). The
- 15 hypocotyl sections were then cocultivated with Agrogribacterium for two days using the same temperature and light conditions used for the seed germination. No feeder layers were used. The cocultivation was terminated by transferring the
- 20 hypocotyl sections to petri plates with liquid callusing medium with 500 mg/l cefotaxime and 200 mg/l vancomycin and gently swirling the plates for about five hours.

The hypocotyl sections were transferred to

- 25 solid callusing medium with 500 mg/l cefotaxime but no selective antibiotics for four days to ensure that the agrobacteria were killed and that the transformed cells could recover from the agrobacterium infection before selection was applied. On the fourth day, the hypocotyl sections were transferred to callusing media with 500 mg/l carbenicillin (Geopen) and 20
 - medie with 500 mg/l carbenicillin (Geopen) and 20 mg/l hygromycin B as the selective antibiotic. The light and temperature regime was the same as that used for seed germination. After 24 days on
- 35 selection, gr en transformed celli could be seen growing from 60% of the cut ends of the hypocotyl

	sections. The negative controls for the
	transformation, consisting of hypocotyl sections not
5	exposed to Agrobacterium, showed no green callus
	growth on media with selective antibiotic.
	After 30 days, the calli were large enough (1
	to 3 mm) to be excised from the hypocotyl sections.
	The excised hypocotyls were transferred to
10	regeneration medium IT-15 (MS salts, B5 vitamins, 3%
	sucrose, 3 mM MES buffer, 2.5 mM IBA, 15 mM Dropp
	(thidiazeron), 0.2% Gel-rite, pH 5.8; supplemented
	with 500 mg/l Geopen, 20 mg/l hygromycin B). This
	medium supports healthy callus growth and rapid
15	regeneration of shoots from non-selected hypocotyl
	sections. The transformed calli are currently being
	tested on this media for rapid organogenesis.
	Plants will be regenerated from calli when
	their diameters have reached at least 0.5 cm by
20	transferring them to KR medium containing 500 mg/l
	Geopen and 20 mg/l hygromycin. KR medium consists of
	K3 major salts (35 mM KNO $_3$, 1 mM (NH $_4$) $_2$ SO $_4$, 1 mM
	${ m MgSO_4}$, 1.5 mM ${ m KH_2PO_4}$, 3.1 mM ${ m NH_4HO_3}$, ${ m CaCl_2}$ added to
	6.3 mM after autoclaving, MS micronutrients, B5
25	vitamins, 1% sucrose, 0.025% xylose, 3 mM MES buffer,
	0.1 mg/l IAA, 2 mg/l zeatin,0.25% low EEO agarose, pH
	5.7). At two week intervals, the outer layers of the
	calli will be trimmed off with a scalpel and they
	will be transferred to fresh media. When shoots have
30	regenerated from the calli, they will be cut away
	from the callus and transferred to Magenta boxes
	containing rooting medium (0.5% MS salts, MS
	micronutrients B5 vitamins, 1% sucrose, 3 mM MES
25	buffer, 0.8% TC agar, pH 5.8) containing 500 mg/1

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boxes will be raised slightly and the opening sealed with Micropore tape to allow ethylene to escape.

The regenerated transformants will be transferred to hydroponics, grown, and treated with N-(aminocarbonyl)-2-chlorobenzenesulfonamide as described in Example 23. It is expected that Brassica plants transformed with the pJE519 construction will show the induction of both GUS mRNA

and GUS enzyme activity upon chemical treatment.

Example 26

Induction of the 2-1, 2-2, and 5-2 corn genes in vivo by Various Chemical Compounds

The ability at various substituted benzenesulfonamides and related compound to induce the expression of the 2-1, 2-2, and 5-2 genes of Missourt 17 corn was evaluated. Corn seeds were

germinated and grown hydroponically in 2 liter beakers as described in Example 1. On the tenth day, plants were transferred into fresh 0.5X Hoagland's solution containing the chemical to be tested. Root tissue was harvested from the plants after six hours of chemical treatment, quick-frozen by immersion in liquid N2, and stored at -80°C until analyzed.

Slot Blot Analysis of RNA from Chemically Treated Corn Plants

Details of the RNA isolation and slot blot analysis procedures are presented in Example 1. Total RNA was prepared from the root tissue of plants that had been treated with various chemicals using the previously described guanidine thiocyanate procedur. Replicate bl ts, each consisting of 2 µg of total RNA from tissues treated with each of the

	chemicals shown in tables 8 and 9, were prepared on
	nitrocellulose membranes using a Minifold II®
5	Slot-blotter (Schleicher & Schuell) following the
	manufacturer's recommeneded procedure. Replicate
	blots were prehybridized and hybridized with comp
	probes made by nick translation of the purified cDNA
	inserts from plasmid pIn 2-1, pIn 2-2-3, and pIn
10	5-2. Slot blots were washed as described in Example
	1 and exposed to Kodak X-OMAT XAR-5 film for 24 hours
	at -80°C using using a single Du Pont Lightning Plus
	intensifying screen. Film was developed using a
	Kodak X-OMAT film processor The Alabama a
15	Kodak X-OMAT film processor. The ability of a
	chemical to induce the mRNA encoded by the three
	inducible genes was evaluated in one of two ways.
	Qualitative evaluation was performed by direct visual
	comparison of the autoradiographic signal intensities
	on the films for the hybridization of each probe to
20	the different RNA samples. Quantitative evaluation
	was performed by cutting each slot containing
	hybridized RNA from the blot, immersing it in 2 ml of
	Du Pont ECONOFLUOR® scintillation cocktail and
	counting the radioactivity in each slot in a
25	scintillation counter. The net amount of
	radioactivity hybridizing to N-(aminocarbonyl)-
	2-chlorobenzenesulfonamide-treated RNA after
	subtraction of radioactivity hybridizing to untreated
	RNA is presented in Table 15.
30	Freedomed In table 15.

TABLE ..

			TABLE 15	1	
	Compound*	In 2-1	In 2-2	In 5-2	
10	1 2 3 4 5 6 7 8	204 111 70 295 296 244 251	332 270 260 237 136 135 129	47 58 61 76 59 53 72 47	
15	10 11 12 13	53 203 102 49 60	110 94 70 8 1	33 63 36 14 55	
	*The n	ames of the	compounds	tested for	4 m.a

of the 2-1, 2-2 and 5-2 promoters in corn roots are

20 listed below. All compounds were used at a concentration of 200 mg/l.

- diethyl [[2-[(butylaminocarbonyl)aminosulfonyl]phenyl]]phosphonate
- 2. N'-[2-(n-butylaminocarbonyl)]-6-chloro-N,N
 - dimethyl-1,2-benzene-disulfonamide
 - 3. N-isopropylcarbamoylbenzenesulfonamide
 - 2-chloro-N-(methylaminocarbonyl)benzenesulfonamide
- 30 5. N-(aminocarbonyl)-2-chlorobenzenesulfonamide
 - 6. 1-cyclohexy1-3-methylsulfonylurea
 - 7. 1-buty1-3-methylsulfonylurea
 8. 2-chloro-N-1/3 /3
 - 2-chloro-N-[[3-(2-ethoxyethoxy)propyl]aminocarbonyl]benzenesulfonamide

TABLE 15 (continued)

- 2,3-dichloro-N-[(cyclopropylamino)carbonyl]benzenesulfonamide
- methyl 2-[(aminocarbonyl)aminosulfonyl]benzoate
- 10 11. N-(aminocarbony1)-2,3-dichlorobenzensulfonamide
 - 2,3-dichloro-N-[(cyclopentylamino)carbonyl]benzenesulfonamide
 - N-(aminocarbonyl)-4-(1,1-dimethylethyl)-2nitrobenzenesulfonamide

15

The responsiveness of the 2-1, 2-2 and 5-2 genes of Missouri 17 corn to hydroponic application of plant hormones and various chemical compounds

associated with plant stress was examined. In

addition, the responsiveness of the corn genes to stress stimuli was also examined. The results are summarized in Table 16.

25

TABLE 16

	5	PLANT HORMONES	In2-1	In2-2	In5-2	
;		Abscisic acid (100 ppm)	+	-	_	
:		6-Benzyladenine (benzyl amino purine) (100 ppm)	++	_	_	
-	10	<pre>2,4-dichlorophenoxyacetic acid (100 ppm)</pre>	+++	+	_	
		Gibberellic acid (100 ppm)	-	_	_	
		Indole acetic acid (100 ppm)	+++	+	n/a	
		Indole butyric acid (100 ppm)	++	+	n/a	
	15	Naphthaleneacetic acid (100 ppm)	+	-	-	
		p-chlorophenoxyacetic acid (100 ppm)	**	++	+	•

20 STRESS STIMULATION

Acetylsalicylic acid (200 ppm)	++	++	++
NaC1 (100 mM)	-	_	_
Proline (20 mM)	-	_	_
Salicylic acid (200 ppm)	+	+	
Salicylamide (200 ppm)	++	·	•
Urea (100 mM)		-	-

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A maximum induction level is represented by

"+++++". This was correlated to the level of induction routinely observed with 100 ppm N-Aminocarbonyl-2-chlorobenzenesulfonamide as the inducing compound.

Example 27

Response of a Recombinant Gene Whose Expression is Controlled by 8 2-2 corn promoter to substituted benzenesulfonamides and Structurally Related Compounds in Transformed Rice Protonlasts

The ability of various substituted benzenesulfonamides and related compounds to induce the expression of recombinant genes consisting of a

GUS coding region driven by regulatory sequence derived from the 2-2 corn gene was evaluated in transformed rice protoplasts. Details concerning establishing rice suspension cultures, isolation and transformation of protoplasts, and the assay of GUS activity were described in Example 14.

Rice protoplasts were transformed with the recombinant DNA construction pTDS133 and then treated with different compounds at a concentration of 100 µg/1 as described in Exemple 10. Table 17 summarizes the results of two such analyses. A

summarizes the results of two such analyses. A number of substituted benzenesulfonamides tested demonstrated the ability to induce GUS activity in transformed protoplasts, with N'-[2-(n-butyl-aminocarbonyl)]-6-chloro-N,N-dimethyl-1,2-benzenedisulfonamide being most active.

In this example, the ability of the various substituted benzenesulfonamides to induce the

- substitute betweensulfonamices to induce the
 expression of a recombinant 2-2 promoter/GUS
 construction in transformed rice protoplasts is shown
 to correlate with the ability of the same compounds
 to induce the expression of the endogenous 2-1 and
 2-2 genes in hydroponically grown Missouri 17 corn
 (Example 17). This indicates that the rice
 protoplast transient assay system is a valuable
 predictive method for determining the ability of a
 chemical to induce genes whose expression is
 regulated by promoters that are inducible by
- substituted bezenesulfonamides and related compounds in whole plants.

TABLE 17

AVE. INDUCTION	
0	
i	
7.67	
29.8	
7.75	
4	
7.15	
20.75	
9.3	
	1.6 3.6 17 16 27.3 24 30 16.6 1.6 5.2 38.6 24.2

	mbt - t - t	
 -	The chemical names of the compounds tested for	
5	induction of the 2-2 promoter/GUS fusion are listed	
3	below:	•
	1. NO DNA	:
	2. 35S-GUS control	
	 methyl 2-[(aminocarbonyl)aminosulfonyl]benzoate 	
10	 N'-butylaminocarbonyl-6-chloro-N,N-dimethyl-1,2- 	
	benzenedisulfonamide	
	N-(aminocarbonyl)-2-chlorobenzenesulfonamide	
	 N-(aminocarbonyl)-4-(1,1-Dimethylethyl)-2-nitro- 	
	benzenesulfonamide	
15	 N-(aminocarbony1)-2,3-dichlorobenzenesulfonamide 	
	 2,3-dichloro-N-[(cyclopentylamino)carbonyl]- 	
	benzenesulfonamide	
 	9. 2-chlore-N-(methylaminocarbonyl)benzene-	
	sulfonamide	
20	10. α -[(1,3-dioxolan-2-yl-methoxy)-imino]-	
	benzeneacetonitrile	
	 phenylmethyl 2-chloro-4-(trifluoromethyl)-5- 	
	thiazolecarboxylate	
	 methyl 3-[(butylaminocarbonyl)-aminosulfonyl]- 	
25	2-thiophenecarboxylate	
	 methyl 2-[[(butylamino)aminosulfonyl]-6-chloro- 	
	benzoate	
	 methyl 3-[(butylaminocarbonyl)aminosulfonyl]-2- 	
	furancarboxylate	
30	15. N-[(butylamino)carbonyl]-3-methyl-2-propyl-	
	sulfonyl-benzenesulfonamide	
	<pre>16. N'-[(butylamino)carbonyl]-N-methyl-N-(1,1,2,2,-</pre>	:
	tetrafluoroethyl)-1,2-benzenedisulfonamide	
	 2-methoxy-6-methyl-N-(methylaminocarbonyl)- 	
35	benzenesulfonamide	

18.	N,N-dimet	ny1-2-[(amin	ocarbonyl)	aminosulfonyl]-3
	Pyridine	carboxamide		

- 5 19. N-(butylaminocarbonyl)-4-chloro-3-pyridinesulfonamide
 - 20. N-(propylaminocarbonyl)-2-pyridinesulfonamide
 - 21. 2,6-dichloro-N-[(1,1-dimethyl)aminocarbonyl]-3pyridinesulfonamide

Example 28

Induction of the Petunia P6 Gene and the Tobacco T2 Gene by Salicylic Acid

- 15 Petunia and tobacco plants were grown as described in Example 5 and treated hydroponically with either 200 mg/l of N-(aminocarbonyl)-2-chlorobenzenesulfonamide or 100 mg/l of salicylic acid for 2, 4, 6 and 22 hours. Total RNA was isolated from
- the roots of treated plants and analyzed for the expression of PG mRNA by RNAse protection as described in Example 4. P6 RNA was detectable by 2 hours following N-(aminocarbonyl)-2-chlorobenzene-sulfonamide treatment and reached maximum levels by 6
- 25 hours. However, maximal levels of P6 RNA were seen by 2 hours following salicylic acid treatment, and this level declined to that seen in untreated plants by 6 hr. This result may suggest a different mode of action for the chemicals.

UTILITY

The promoters shown in figures 2, 4, 5 and 7 are useful for regulating the expression of 5 structural genes operably linked to plant promoters derived from the genes in response to the external application of compounds of the Formulae I-IX. Regulation of genes is achieved by application of the 10 compounds of formulae I-IX to transgenic plants containing chimeric genes consisting of structural genes encoding a gene product to be regulated operably linked to promoters described in figures XX-YY and their derivatives. 15 A number of methods are available for application of the inducing compounds described herein. The inducer may be applied directly to the

crop seed. The seeds may be uniformly coated with the inducer according to standard seed treating procedures prior to planting. Alternatively, the

inducer may be applied over the the exposed seeds in open furrows at planting, just prior to covering the seed with soil (in-the-furrow treatment). The inducer may be applied post-emergence at the specific 25 time that expression of the desired gene(s) is

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appropriate. Post emergent application may be directed so that the inducer is primarily applied to the crop. The amount of inducer will vary depending on the specific inducer and the method of application

30 used. The crop species and cultural practices may also have an effect.

It is expected that regulating the temporal expression of genes responsible for a number of plant traits will be agronomically beneficial in transgenic plants. Examples of traits include herbicide resistance where limiting a plant's resistance to a

class of herbicide(s) by controlling the expression of a gene conferring herbicide resistance would be beneficial. In this manner, unwanted volunteer plants germinating in the field as a result of seed lost during the harvest could be easily eliminated if the inducing gene were left unactivated. Examples of such herbicide resistance genes include resistant forms of the acetolactate synthase gene (sulfonylurea 10 herbicide resistance), the 5-enolpyruvylshikimate-3-phosphate synthase gene (glyphosata resistance), and the BAR gene (encoding Basta resistance). 15 Controlling the expression of genes conferring pathogen and insect resistance would also be of agronomic benefit. By limiting the expression of these resistance genes to the times in the pest's life cycle when infestation occurs, one would limit the rate of appearance of resistance to the gene 20 product in the pest population by limiting the expression of the resistance genes to short periods of time. Restricting the expression of resistance genes to relatively short times during the growth 25 cycle of the plant may well minimize any yield penalty associated with constitutive expression of the desired gene. Examples of such genes include any of genes encoding Bacillus thurengensis insecticidal endotoxins, chitinase genes, protease inhibitor 30 genes, genes encoding nematode resistance and so on. In addition by using recombinant, chemically inducible promoters one may be able to express a pest toxin in only affected tissues and prevent their expression in portion(s) of the plant to be used as 35 foodstuffs.

	enemically regulating the expression of genes
	involved in pytchormone biosynthesis in transgenic
5	plants may have agricultural benefit. For example,
	chemical induction of 1-amino-cyclopropane-1-
	carboxylic acid synthase genes just prior to harvest
	may accelerate fruit ripening as a harvest aid by
	providing a burst of ethylene synthesis immediately
10	prior to harvest. Similarly, regulating the
	expression of other genes involved in the
	biosynthesis of other phytohormones such as
	cytokinins, auxins, gibberellins, and abscisic acid
	to control hormone levels in field grown plants may
15	prove to have great agricultural utility.
	There would be substantial agronomic benefit in
	regulating the expression of a great number of plant
	traits if one knew which gene(s) encode the
	protein(s) responsible for these traits. As these
20	
20	genes and their products are discovered, regulating
	their expression by external chemical control may
	well have agronomic value. In this manner, yield
	penalties associated with constitutive expression of
	a trait that may be needed for a relatively short
25	period of time, can be minimized. Examples of such
	genes and traits are drought resistance genes, salt
	tolerance genes, pathogen resistance genes, and so on.
	By expressing genes for degradative enzymes in
	specific plants tissues just prior to harvest, one
30	may be able to reduce the processing costs associate

costs for the brewing industry, increasing the yield 35 of sucrose in sugarbeets by expression of just prior to harvest, improving the nutritional

with converting raw plant materials to useable forms. Examples include the expression c-amylase in rice seeds just prior to harvest to reduce processing

· :	5	quality of soybeans by reducing raffinose and raffinosaccharides in by expression of high levels of $\alpha\text{-galactosidase}$ in seeds just prior to harvest, expression of ligninase in plant tissues used by the pulp and paper industries.
	10	•
	15	
	20	<u>. </u>

CLAIMS
What is claimed is:

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 A nucleic acid promoter fragment inducible by a compound of Formula I-IX:

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VI

15 N BO₂NHCONHR

wherein

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X is H, F, Cl, Br CF₃, or C_1 - C_2 alkyl; Xl is H, F, Cl, C_1 - C_2 alkyl, $SO_2NR^1R^2$ or CO_2R^1 ; Y is H, Cl or $SO_2NR^1R^2$, CO_2R^1 , NO_2 , $P(O)(OR^1)_2$; R is H, C_1 - C_3 alkyl, C_3 - C_6 evoloalkyl, benzyl or C_2 - C_4 haloalkyl or C_2 - C_4 substituted with C_1 - C_2 alkoyr or C_1 - C_2 alkylthio; Rl is C_1 - C_3 alkyl; R2 is C_1 - C_3 alkyl; R2 is C_1 - C_3 alkyl;

 R^3 is CO_2R_2 ; R^4 is C_1-C_6 alkyl or C_3-C_6 cycloalkyl;

20

R ⁵	is C ₁ -C ₃ alkoxy or NR ⁶ R ⁷ ;
R	is H, OCH ₃ , C_1 - C_4 alkyl, C_3 - C_6 cycloalkyl, C_1 - C_4 alkyl substituted with C_1 - C_2 alkoxy o
	ethoxyethoxy; and

R⁷ is H or C₁-C₂ alkyl; and agriculturally suitable sa

and agriculturally suitable salts thereof such that exposure of plants transformed with said promoter

- fragment to a compound of Formula I-IX causes increased expression of a DNA sequence coding for a selected gene product operably linked to said promoter fragment.
- 2. A nucleic acid promoter fragment of Claim 1 wherein the compound of Formula I is a member of the group wherein:

X is H or 2-Cl;

Y is 3-Cl or $SO_2N(CH_3)_2$;

R is H, C₁-C₆ alkyl or C₅-C₆ cycloalkyl.

3. A nucleic acid promoter fragment of Claim 1 wherein the compound of Formula II is a member of the group wherein:

25 R is C_1 - C_4 alkyl or C_5 - CX_6 cycloalkyl; R_4 is C_1 - C_4 alkyl.

 A nucleic acid promoter fragment of Claim 1 wherein the compound of Formula III is a 30 member o the group wherein:

> R_5 is OCH₃ or NR₆R₇; R_6 is H or C₁-C₄ alkyl; and R_7 is H.

35 5. A nucleic acid promoter fragment of Claim 1 wherein said compound of Formula I-IX is a compound selected from the group consisting of

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diethy1[[2-(butylaminocarbonyl)aminosulfonyl]phenyl]]phosphate, N-isopropylcarbamoylbenzenesulfonamide, 1-cyclohexyl-3-(methylsulfonyl)ures,
1-(n-butyl)-3-methylsulfonylures, methyl-2[(aminocarbonyl)aminosulfonyl]benzoate,
N-(aminocarbonyl)-2-chlorobenzenesulfonamide,
N'-[2-(n-butylaminocarbonyl)]-6-chloro-N,N-dimethyl1,2-benzenedisulfonamide, 2-chloro-N-(methylamino-

- carbonyl)benzenesulfonamide, 2,3-dichloro-N-[(cyclopentylamino)carbonyl)]benzenesulfonamide, and N-(aminocarbonyl)-2,3-dichlorobenzenesulfonamide.
- 15 6. A nucleic acid promoter fragment of Claim 1 derived from a plant.
 - 7. A nucleic acid promoter fragment of Claim 6 wherein said plant is a monocotyledonous plant.
 - 8. A nucleic acid promoter fragment of Claim 7 wherein said monocotyledonous plant is selected from the group consisting of corn, oats, millet, wheat, rice, barley, sorghum, amaranth, onion, asparagus and sugar cane.
- A nucleic acid promoter fragment of Claim
 8 wherein said monocotyledonous plant is selected
 from the group consisting of corn and rice.
 - A nucleic acid promoter fragment of
 Claim 6 wherein said plant is a dicotyledonous plant.
- 35 11. A nucleic acid promoter fragment of Claim 10 wherein said plant is a dicotyledonous plant selected from the gr up c nsisting of alfalfa, soybean, petunia, cotton, sugarbeet, sunflower, carrot, celery, cabbage, cucumber, pepper, canola,

tomato, potato, lentil, flax, broccoli, tobacco, hean, lettuce, oilseed rape, cauliflower, spinach, brussel sprout, artichoke, pea, okra, squash, kale,

- 5 brussel sprout, artichoke, pea, okra, squash, kale, collard greens, tea and coffee.
- 12. A nucleic acid promoter fragment comprising a nucleotide sequence from the 5' flanking promoter region of a corn gene substantially homologous to cDNA clone 2-1 deposited with the American Type Culture Collection (ATCC) and given the ATCC accession designation 67805.
- 15 13. A nucleic acid promoter fragment of Claim 12 comprising a nucleotide sequence from the 5' flanking promoter region of a corn gene substantially homologous to CDNA clone 2-1 deposited with the
- American Type Culture Collection (ATCC) and given the 20 ATCC accession designation 67805, such that exposure of plants transformed with said promoter fragment to a compound of Formula I-IX causes increased expression of a DNA sequence coding for a selected gene product operably linked 3' to said promoter fragment.
 - 14. A nucleic acid promoter fragment of Claim 13 wherein said compound of formulae I-IX is compound selected from the group consisting of N-(aminocarbonyl)-2-chlorobenzenesulfonamide, 2-chloro-N-(methylaminocarbonyl)benezenesulfonamide, 1-(n-butyl)-3-methylsulfonylurea, 1-cyclohexyl-3-(methylsulfonylurea, diethyl [[2-(butylaminocarbonyl)aminosulfonyl)phonyl]] phosphonate, methyl
- 35 2-[(aminocarbonyl)aminosulfonyl]benzoate, 2,3-dichloro-N-[(cyclopentylamino)carbonyl]benzene-

sulfonamide, and N-(aminocarbonyl)-2,3-dichlorobenzenesulfonamide.

- 15. A nucleic acid promoter fragment of Claim 14 wherein said compound of Formula I-IX is N-(aminocarbonyl)-2-chlorobenzene sulfonamide.
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 16. A nucleic acid promoter fragment of Claim 11 comprising the nucleotide sequence of 590 base pairs running in the 5' to 3' direction from base pair position 1 to base pair position 590 as shown in Figure 2 from a gene substantially

 15 homologous to CDNA clone 2-1.
 - 17. A nucleic acid promoter fragment of Claim 16 comprising the nucleotide sequence of 363 base pairs corresponding to base pairs 169 to 532 in
- 20 Figure 2, or any promoter fragment substantially homologous therewith.
- 18. A nucleic acid acid promoter fragment comprising a nucleotide sequence from the 5' flanking promoter region of a corn gene substantially homologous to cDNA clone 2-2 deposited with the American Type Culture Collection (ATCC) and given the ATCC accession designation 67803
- 30 19. A nucleic acid promoter fragment of Claim 18 comprising a nucleotide sequence from the 5filanking promoter region of a corn gene substantially homologous to cDNA clone 2-2 deposited with the American Type Culture Collection (ATCC) and given the
- 35 ATCC accession designation 67803, such that exposure

of plants transformed with said promoter fragment to a compound of Formula I-IX causes increased

- expression of a DNA sequence coding for a selected gene product operably linked 3' to said promoter fragment.
- 20. A nucleic acid promoter fragment of Claim 19 wherein said compound of formulae I-IX is a 10 compound selected from the group consisting of diethyl [[2-(butylaminocarbonyl)aminosulfonyl]phenyl] phosphonate, N'-[2-(n-butylaminocarbonyl)]-6chloro-N,N-dimethy1-1,2-benzenedisulfonamide.
- N-isopropylcarbamoylbenzenesulfonamide, 2-chloro-N-15 (methylaminocarbonyl)benzenesulfonamide, N-(aminocarbonyl)-2-chlorobenzenesulfonamide, and 1-cyclohexy1-3-(methylsulfonyl)urea.
- 20 A nucleic acid promoter fragment of 21 Claim 20 wherein said compound of Formula I-IX is diethyl[[2-(butylaminocarbonyl)aminosulfonyl]phenyl] phosphonate.
- 25 22. A nucleic acid promoter fragment of Claim 18 comprising the nucleotide sequence of 207 base pairs running in the 5' to 3' direction from base pair position 264 to base pair position 470 as shown in Figure 4 from a gene substantially homologous to cDNA clone 2-2. 30
 - 23. A nucleic acid promoter fragment of Claim 22 comprising the nucleotide sequence of 77 base pairs corresponding to base pairs 292 to 368 in
- Figure 4, or any promoter fragment substantially homologous therewith.

A nucleic acid promoter fragment

comprising a nucleotide sequence from the 5' flanking promoter region of a corn gene substantially homologous to cDNA clone 5-2 deposited with the American Type Culture Collection (ATCC) and given the ATCC accession designation 67804.

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- 25. A nucleic acid promoter fragment of Claim 24 comprising a nucleotide sequence from the 5' flanking promoter region of a corn gene substantially homologous to cDNA clone 5-2 deposited with the American Type Culture Collection (ATCC) and given the ATCC accession designation 67804, such that exposure of plants transformed with said promoter fragment to a compound of Formula I-IX causes increased
- expression of a DNA sequence coding for a selected
 gene product operably linked 3' to said promoter
 fragment.
- 26. A nucleic acid promoter fragment of Claim 25 wherein said compound of formulae I-IX is a compound selected from the group consisting of 2-chloro-N-(methylaminocarbonyl) benzenesulfonamide, 1-(n-butyl)-3-methylsulfonylurea, methyl 2-[(aminocarbonyl)aminosulfonyl]benzoate, N-isopropylcarbamoyl-benzenesulfonamide, N-(aminocarbonyl)-2-chlorobenzenesulfonamide and N'-[2-(n-butylaminocarbonyl)]-6-chloro-N,N-dimethyl-1,2-benzenedisulfonamide
 - 27. A nucleic acid promoter fragment of Claim 26 wherein said compound of Formula I-IX is 2-chloro-N-(methylaminocarbonyl)benzenesulfonamide.

- 28. A nucleic acid promoter fragment of Claim 24 comprising the nucleotide sequence of 889
- 5 base pairs running in the 5' to 3' direction from base pair position 1 to base pair position 889 as shown in Figure 5 from a gene substantially homologous to cDNA clone 5-2.
- 10 29. A nucleic acid promoter fragment comprising a nucleotide sequence from the 5' flanking promoter region of a petunia gene substantially homologous to cDNA clone P6.1 deposited with the American Type Culture Collection (ATCC) and given the 15 ATCC accession designation 67823.
 - 30. A nucleic acid promoter fragment of Claim 29 comprising a nucleotide sequence from the 5° flanking promoter region of a petunia gene
- substantially homologous to cDNA clone P6.1 deposited with the American Type Culture Collection (ATCC) and given the ATCC accession designation 67823, such that exposure of plants transformed with said promoter fragment to a compound of Formula I-IX causes
- 25 increased expression of a DNA sequence coding for a selected gene product operably linked 3' to said promoter fragment.
- 31. A nucleic acid promoter fragment of
 30 Claim 30 wherein said compound of Formula I-IX is a
 compound selected from the group consisting of
 N-(aminocarbonyl)-2-chlorobenzenesulfonamide,
 N'-[2-(n-butylaminocarbonyl)]-6-chloro-N,N-dimethyl1,2-benzenedisulfonamide, 2-chloro-N-(methylamino-

carbonyl)benzenesulfonamide, 2,3-dichloro-N-[(cyclo-pentylamino)carbonyl)]benzenesulfonamide, and

N-(aminocarbony1)-2,3-dichlorobenzenesulfonamide.

32. A nucleic acid promoter fragment of Claim 31 wherein said compound of Formula I-IX is N-(aminocarbonyl)-2-chlorobenzenesulfonamide.

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- 33. A nucleic acid promoter fragment of Claim 29 comprising the nucleotide sequence of 595 base pairs running in the 5' to 3' direction from base pair position 1 to base pair position 595 as shown in Figure 8 from a gene substantially homologous to CDNA clone ps. 1.
- 34. A nucleic scid promoter fragment of

 Claim 33 comprising the nucleotide sequence of 240
 base pairs corresponding to base pairs 356 to 595 in
 Figure 8, or any promoter fragment substantially
 homologous therewith.
- 35. A nucleic acid promoter fragment
 25 comprising a nucleotide sequence from the 5' flanking
 promoter region of a tobacco gene substantially
 homologous to cDNA clone T-2.1 deposited with the
 American Type Culture Collection (ATCC) and given the
 ATCC accession designation 67822.

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36. A nucleic acid promoter fragment of Claim 34 comprising a nucleotide sequence from the 5° flanking promoter region of a tobacco gene substantially homologous to cDNA clone T2.1 deposited with th American Type Culture Collection (ATCC) and

given the ATCC accession designation 67822, such that exposure of plants transformed with said promoter

- fragment to a compound of Formula I-IX causes increased expression of a DNA sequence coding for a selected gene product operably linked 3' to said promoter fragment.
- 10 37. A nucleic acid promoter fragment of Claim 36 wherein said compound of Formula I-IX is a compound selected from the group consisting of N-(aminocarbony1)-2-chlorobenzenesulfonemide, N'-[2-(n-butylaminocarbony1)]-6-chloro-N,N-dimethyl-
- 15 1,2-benzenedisulfonamide, 2-chloro-N-(methylamino-carbonyl)benzenesulfonamide, 2,3-dichloro-N-[(cyclo-pentylamino)carbonyl)]benzenesulfonamide, and N-(aminocarbonyl)-2,3-dichlorobenzenesulfonamide.
- 38. A nucleic acid promoter fragment of Claim 37 wherein said compound of Formula I-IX is N-(aminocarbonyl)-2-chlorobenzenesulfonamide.
- 39. A nucleic acid promoter fragment 25 comprising a nucleotide sequence from the 5'flanking promoter region of a corn gene substantially homologous to cDNA clone 218.
- 40. A nucleic acid promoter fragment of Claim
 30 39 comprising a nucleotide sequence from the 5'
 flanking promoter region of a corn gene substantially
 homologous to cDNA clone 218, such that exposure of
 plants transformed with said promoter fragment to a
 compound of Formula I-IX causes increased expression
 35 of NNA sequence coding for sealers.
- 35 of DNA sequence coding for a selected gene product operably linked 3' to said promoter fragment.

- A nucleic acid promoter fragment of
- 5 Claim 40 wherein said compound of formulae I-IX is compound selected from the group consisting of N-(aminocarbonyl)-2-chlorobenzenesulfonamide, 2-chloro-N-(methylaminocarbonyl)benzenesulfonamide, 1-(n-butyl)-3-methylsulfonylurea, 1-cyclohexyl-3-10 (methylsulfonyl)urea, diethyl [[2-(butylamino-carbonyl)aminosulfonyl]phosphonate, methyl 2-[(aminocarbonyl)aminosulfonyl]benzoate, 2,3-dichloro-N-[(cyclopentylamino)carbonyl]benzenesulfonamide, and N-(aminocarbonyl)-2,3-dichloro-benzesulfonamide
 - 42. A nucleic acid promoter fragment of Claim 41 wherein said compound of Formula I-IX is N-(aminocarbony1)-2-chlorobenzene sulfonamide.

- 43. A nucleic acid promoter fragment of Claim 39 comprising the nucleotide sequence of 1574 base pairs running in the 5' to 3' direction from base pair position 1 to base pair position 1574 as shown in Figure 7 from a gene substantially homologous to cDNA clone 218.
- 44. A recombinant DNA construct, capable of transforming a plant, comprising a nucleic acid promoter fragment of Claims 1-42 or 43, a DNA sequence coding for a selected gene product operably linked to said promoter fragment, and a suitable 3' downstream region such that exposure of said transformed plant to a compound of Formula I-IX auses increased expression of said DNA sequence for a selected gene product.

45.	Α	recombinant	DNA	construct	of	Claim 44

- wherein said DNA sequence for a selected gene product is selected from the group consisting of the sequence for β-glucuronidase, acetolactate synthase, 5-enolpyruvylskikimate-3-phosphate synthase, a gene encoding a product capable of conferring insect resistance, a gene encoding a protease inhibitor, a
- gene encoding a Bacillus thuringiensis insecticidal endotoxin, a gene encoding phytohormone biosynthesis, a gene encoding l-amino-cyclopropane-l-carboxylic acid synthase, a gene encoding auxin biosynthesis, a gene encoding cytokinin biosynthesis, a gene encoding cytokinin biosynthesis, a gene encoding cytokinin biosynthesis, a gene encoding
- 15 giberellin biosynthesis, a gene encoding chitinase, and a gene encoding biosynthetic enzymes for oil production.
- 46. A recombinant DNA construct, capable of transforming a plant, comprising (1) a nucleic acid promoter fragment of Claim 23 inserted into (2) a promoter sequence selected from the group consisting of the CaMV 19S and 3SS promoters, and NOS and OCS promoters of the opine synthase gene of
- 25 Agrobacterium, the promoter of the small subunit of RUBISCO, the promoter from the chlorophyll A/B binding protein genes, a root specific promoter, a leaf specific promoter, a stem specific promoter, a seed specific promoter, a pollen specific promoter,
- 30 an ovule specific promoter, a stress-inducible promoter, a developmentally regulated promoter, and a constitutive promoter, (3) a DNA sequence coding for a selected gene product operably linked to said promoter sequence, and (4) a suitable 3' downstream
- 35 region such that said promoter sequence causes increased expression of said DNA sequence for a selected gene product upon exposure of said transformed plant to a compound of Formula I-IX.

- 47. A transgenic plant containing a nucleic acid promoter fragment of Claims 1-42 or 43 such that exposure of said transgenic plant to a compound of Formula I-IX causes increased expression of a DNA sequence coding for a selected gene product operably linked 3° to said promoter fragment.
- 10 48. A transgenic plant of Claim 47 wherein said plant is a monocotyledonous plant selected from the group consisting of corn, oats, millet, wheat, rice, barley, sorghum, emaranth, onion, asparagus and sugar cane.
 - 49. A transgenic plant of Claim 48 wherein said plant is a monocotyledonous plant selected from the group consisting of corn and rice.
- 50. A transgenic plant of Claim 47 wherein said plant is a dicotyledonous plant selected from the group consisting of alfalfa, soybean, petunia, cotton, sugarbeet, sunflower, carrot, celery, cabbage, cucumber, pepper, canola, tomato, potato, lentil, flax, broccoli, tobacco, bean, lettuce,
- oilseed rape, cauliflower, spinach, brussel sprout, artichoke, pea, okra, squash, kale, collard greens, tea, coffee, geranium, carnation, orchid, rose, impatiens, petunia, begonia, fuscia, marigold, chrysanthemum, cladical
 - 0 chrysanthemum, gladiola, astromeria, salvia, veronica, daisey, and iris.
 - Seed obtained by growing a transgenic plant of Claim 47.

 A method of causing increased expression
of a selected gene product in a plant comprising the
steps of (a) transforming said plant with a
recombinant DNA construct of Claim 44, (b) exposing
said transgenic plant to a compound of Formula I-IX,
and (c) causing said transgenic plant to increase

- said transgenic plant to a compound of Formula I-IX, and (c) causing said transgenic plant to increase expression of said selected gene product at a desired time.
 - 53. A method of causing increased expression of a selected gene product in a dicotyledonous plant comprising the steps of (a) transforming said dicotyledonous plant with a recombinant DNA construct containing a nucleic acid promoter fragment of Claim 29 or 35, (b) exposing said transgenic dicotyledonous plant to salicylic acid, and (c) causing said
- transgenic dicotyledonous plant to increase
 20 expression of said selected gene product at a desired
 time.

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FIG 1

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1. GROW CORN SEEDLINGS HYDROPONICALLY

- ADD N-(AMINOCARBONYL)-2-CHLOROBENZENESULFONAMIDE TO HYDROPONIC MEDIUM OF HALF OF THE PLANTS AND GROW FOR SIX HOURS
- 3. ISOLATE MRNA FROM ROOTS OF TREATED AND UNTREATED PLANTS
- 4. CREATE CDNA LIBRARY FROM MRNA FROM TREATED PLANTS AND REPARE REPLICA COPIES OF LIBRARY
- 5. SCREEN COPIES OF cDNA LIBRARY WITH 32P-DNA PROBES MADE FROM EITHER TREATED OR UNTREATED ROOT MRNA TO ISOLATE CLONE CONTAINING SEQUENCES INDUCED BY N-{AMINOCARBONYL}-2-CHLOROGENZENESULFOVAMIDE
- 6. PREPARE CORN GENOMIC LIBRARY
- 7. USE cDNA CLONE TO ISOLATE CORRESPONDING CHEMICALLY INDUCED GENE(S)
- 8. DETERMINE SEQUENCES OF cDNA CLONE AND GENE. IDENTIFY PROMOTER AND 3' DOWNSTREAM REGIONS OF GENE TO BE REMOVED FROM STRUCTURAL PORTION OF GENE
- 9. ADD CONEVIENENT RESTRICTION SITES FOR CLONING (IF NEEDED) AND CREATE RECOMBINANT GENE BY OPERABLY LINK B-GLUCURONIDASE CODING REGION TO PROMOTER AND 3' DOWNSTREAM REGION OF INDUCIBLE GENE
- 10. TRANSFORM RECOMBINANT GENE INTO PLANTS
- 11. TEST PLANTS FOR N-(AMINOCARBONYL)-2-CHLOROBENZENESULFONAMIDE INDUCIBLE EXPRESSION OF RECOMBINANT GENE

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1	CTACCTTCAT	GAGACGTAAC	TCCLCLL		MINICGGITA
51	TCTT 1 00000		1 GCAGAAGAT	GTECTTTCCA	AITHCGGTTA
	IGITACCTTT	AATCCCAAGC	CTTCAGCELT	GENERICIAT	GETTAACTT
101	CTTATTGAAG	CCAAGATATC	TOTAL COLD		
151	GAGAAAATAG	CACCCON			MERITACGA TINGTAGACAT
201	Champana	- COCINIGE	GCCTTTCTAR	THA GASKICE	TOGTAGACAT

- 201 GACTICAGCA GITTAGGICA TAGATGACGA GITAGGICAGA GGALLIGGCAA
 251 TGGGGCCAAC ACGAATIGIT CGTGCGTCAE ANGURGGGGA ANGUGAACACA
- 301 ATCGATTACG TCATCAGTCG TTTAACTCAA (CUGINACAC) AUGGEGGTCCT
- 351 GACAGGTGGG GCGCCACCGC AATTTATTET TANKCAGCGA GETABECGGCG
- 401 ACAGACACGT GGTGGGCCTG TGGGGGTCTE GTGGCTTRAA CETTERAANG
- 451 TCATGCATGC ACTGCGCTAA AGTCTAAGCC ATLATMAAA UACTGCGCGT
 501 ATAAATACCC GGACCAATCA GCCATGCGGG CAGCCGGGGTC GGGGTTCGAA
- 551 CAGGCCAGTC CCCTCCCACT CCCAGTCCD2 TCTCGACGAY ATTEC

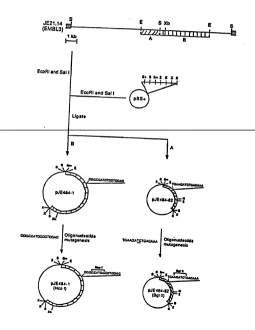
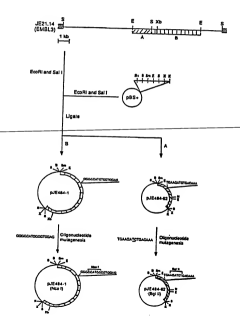


FIG. 3

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PCT/US90/01210



E≈Eco RI S≈Sel I

451 CCCGACTCCC TGCACCTGCC ATGG

AGGARTICET CICCATGGAT CCCCTCTATT TACCTGGCCA CCARACATCC

51 CTAATCATCC CCARATITTA TAGGARCTAC TAATTICTCT AACTTAANAA

101 AAATCTAANA TAGTATACTT TAGCAGCCTC TCAATCTGAT TIGTICCCCA

151 AATTTGAATC CTGGCTTCGC TCTGTCACCT GTTGTACTCT ACATGGTGCG

201 CAGGGGGAGA GCCTAATCTT TCACGACTTT GTTTGTAACT GTTAGCCAGA

251 CCGGCGTATT TGTCAATGTA TAAACACGTA ATAAANTTTA CGTACCATAT

301 AGTAAGACTT TGTATATAAG ACGTCACCTC TTACGTGCAT GGTTATATGC

351 GACATGTGCA GTGACGTTAT CAGGARCTCAT CAGCACCCCG GCAGTGCCAC

FIG. 5

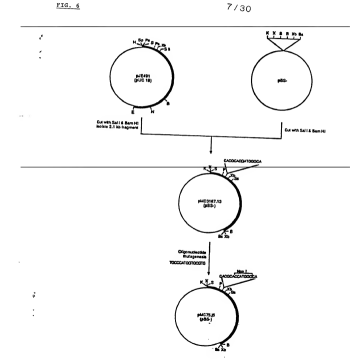
PCT/US90/01210

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1	GCGGTCACAA	TTACCCTATA	TATCTACTAT	ATACCAACTA	CCATTTATTA	
51	TATCATATTT	TTACCATACT	CTATACCAAC	TCCATCACAC	GGCTGCTGTA	
101	CTGCTTCCTT	CTACTGCTAC	TGTACTGGTT	CTCTAGGCCC	ACCTCGTCTG	
151	CTGGGAGAGA	GCAGTGGCAG	AGCGCTACAT	TTGGCGTAGA	AGAGGCGGAG	
201	AGAGAGCGTA	GAGTGAGATA	TAGAGTGCAC	CGTTGCAGAT	CTTGTCTACT	
251	GTAAAANTTT	AGCGTAGCTT	TTCCAGCTGA	CCACTGCGGC	TAGCCTAAAA	
301	CGGATTGGGG	GTACTCAGTG	GNNNNGCCGT	GGGCGGTACG	TCGCCCCAAA	
351	TAATTAAACG	GTGCTCGATG	TACCTCTACG	GGACCTTTTT	CAGCCTTTTT	
401	TCTTTATTTT	ATTATTATTA	TTTTGGTACT	ACACAAGGGA	CCTTTTGACG	
451	CTGAGATGAT	GCCCAAAAAC	AAAAGGACGC	TCATCATCAG	TGACGCCCAG	
501	TCGTCGCCAA	GCAGCTAGCT	AGCATGCCAA	TAATTTTTTT	CTTGTTAATG	
551	TTGTCGCAGC	TGGTACTATA	CTACTACTAC	TACGCCGTAT	ATGAATGCGC	
601	GTTTTGTCTG	ATGCTCAGGC	TGATTCCATC	CAATTGTCTT	TCTTCTCTCC	
651	TCTCCACCCA	TGCCCCGTCC	GTCGCAGCAG	GGGTTATATA	GTGCCCGCGA	

701 ACGGACGCAG GCGCCACGAA GCCGAGATCG AGCAGCTACC TCTCCGATCC
751 GAGGCCTGAG CGAGCGAGCT GAGGACTGCA GCCTATATAA TATCTAGACT
801 AGAGTACACC ACAACGACGA GGCACATATA TATACACGCG GCGGCGCCA
851 GATCCATCTT GGTATACACG TAATATATAT ACACGCACGA TGG

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		WO 90/1	FIG. 7	8/30 PCT/US90/01210
			1 GAATTCGTTT ATAAAAAT	AT ATCGTTCCGC AGGCGTTGAG CCTTTTTCTA
		5	1 CTAGTGATGT CTTCACAAC	OG TTTCGAGCTT TTCCCTAATT GGCGGGTCAT
_		10	1 TAAGGCTTGT ACACGGAGT	C TITCTCCTAC TCTACCCCTG TTAGAAGGCG
		15	1 TAACCCCTTT TTATAAGCC	C GAACACCTGA TGACCAAACC AGGCCAAAGG
		20	1 GTATAACGAT TGTTGCCCC	C CTAATCAGCG CAATAATGCG CGTGGGCCTA
	*	25	ACGCTGTTAA GACTCGATC	C TATTGACCCG TCCGAGATCA ACCTAACAAA
		30:	GTTCTAGCCA TGTGCCATT	T CGTAATGAAA ATGAGGGCCA AGGTGTCACC
		35:	TTGCTGGTCT AAAAAATGT	G CCTCGATCCA AGGGACTGTT CATTTTTAA
		401	AATGACCATA TGACAGACA	T CAGGCTAATG GACATGGTTG AGTTTGGATT
		451	GGCTCAACTC GGTTCGTTA	A CAAACCAATC CAAAAAGTCA GCTCGCTATT
		501	TACGAGCTCG AACAATTATT	F ATCATTAATC AATTTGCTTG TTAGTTACAA
		551	ATTCAGTTTT ACTTAACAGA	A AAAATAGTTA ATTTATTCTT CATAATTTCA
		601	CAGACCATTA TAAATTAAAC	CACTARATTAR TATAGRATCA ATCACAGACA
_		651	TAATTTATCA TCATCAGTTT	GAATCCACGA GCTACATAAG CCGCACATAC
		701	AATGTAGCAT ATTCACCGAT	TCTAGATGAA ATATACTGCA TATAGTTTTA
6		751	TTTTTTGAAN GTGATAGGTC	GTTTGACATC ACGAACTGGC TCGTTAACAA
		801	ACAAGCTAGG ATGTTAGCTT	ATGCTTTGCT ATTAGTTAGG ATATGGTTCT
		851	GGGTGATCAA AAGGAAGAA	AAACACGAAA AATTTAATGA GGTTCTTGGA
		901	TGACCGGAGT CAACCAACTT	AAACACGAAA AATTTAATGA GGTTCTTGGA
		951	GTAGTEGGEA CTETTEGGET	GGTTGGAGCG TTCTTCTTCC CTGATCGTTC
		1001	CGCGAACCAG ACCMAGCOMA	ACGGCTGACG TCCTCACCTC TCCTCGTCCA
		1051	TOTAL MEGIACOGIA	GCTGTTTCAC ATTTCTAATT TACTATACGT
		1101	ACCCTCTING CHECKING	ACCACCTCTC GCATTGCTAA TTTACTGGAT
	:	1151	AAAGTTTATT TTTTGGACACA	AATTGGACCT GCAACGGACT GATGAATTGC
		1201		GAAGGTAAAG CTGAAACGAG TTCCTCCGTC
	•		AAACCTOATA TATTTTGAA	CCGCGAGAGT TCAAATCCCC AGCCAAGCTG
		1201	COCCUCAGA GCCTGAAATT	TTCGTGCTGG GATGACGTTC GCCCTTACGT
•		1301	CGCGCGCTGC AAACTGAAAC	GAGTTCCCAT GCCCARATAA ACTTGAGAAA

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1351 AGTGCTGTCT TGTTCAGCTA TGCCCGCATT ATAGATCGAT ATGGTGAGGT

1401 CACTGCTTAT GCCAGGCACA TGACTCAATA TAGGTCCATA TCTTAGGCGA

1451 ATTAATCACA TCTCTCTGAC CGATCTTGGG GTCCCCTATA AATTATATAGG

1501 AACGTACGTA AAGTTTCTCC AAGCAGATAG CAGCAAGCTA AGCAAGTGCC

1551 AACCAACGAG TAGCAGGAAA CATG

FIG. 8

651 CATAACAATG

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1	ACTGAAGAAT	GATGAGTGAC	TCACAAAATG	GTTTCCCATT	GTCCATCAAC	
51					GAGGACTITT	
101	GATCCTCAAG	TCCTCCTTCC	TIGIATICAT	A A T C A A T T T O	BAGGACTTTT	
151	AGGGCAATCC	TGACCCTCAT	CCCAAACAMA	OMORAL ICC	TITTTCAGCC	
201	AATTTCATCT	GCCTTTTTTT	TTARAMERA	CIGTAAGTAT	CTAGTAGGAC	
251	AATTCCAACA	AATATAAAAC	TIANANIGAA	ATTTAAGGAT	AGTATAATGG	
301	TGAAGTACCA	AATTTCTCCC	TAGAATCAGT	TATTATTCAA	CATAAACCCA	
251	TGAAGTACCA	ANTITIGIGG	GGTAGAGAGA	AGATTTGGAT	CGACTAAAAT	
401	TTTGACTAGT	AAGTTAAAAA .	AATTAAGGAA	Cagaagaaag	TGGAGCCTTC	
401	TTGCTTAACG	TTTACTACTA	TAAGACCCCG	TGACGAATGT	GATGACATAA	
451	GTAGGTCGGC	САСАСААААА :	AATCTGGAAA	CTCCCGGACC	ACAACACCGC	

501 TIGENACCAT ANTANANAG TITANANAG ANGACATCHA AGITTCHACE
551 GGTCTATATA TAGAACTIGA ACTATATACG ANGCATATCA GTTCHAGCA
601 TITGTGCAAA TICTATANAT TCTTCTTACT TGCCTTTCAT AATTCATANAG

FIG. 9

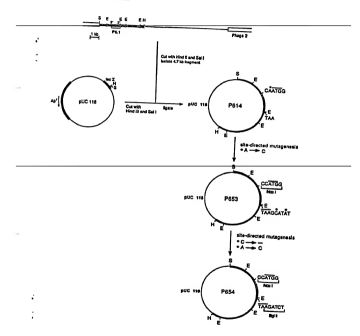
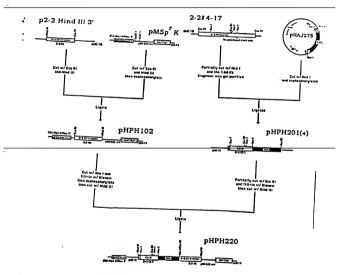
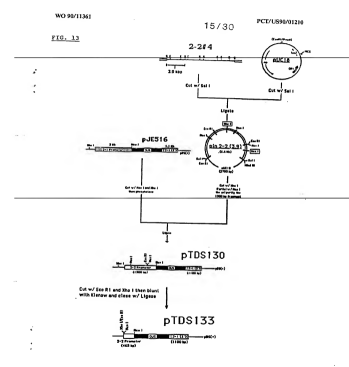


FIG. 12

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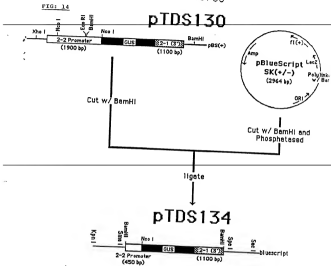




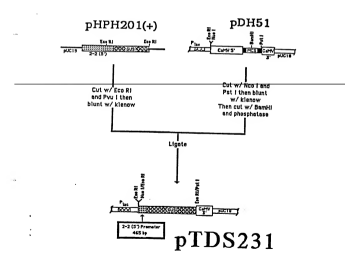


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FIG. 16

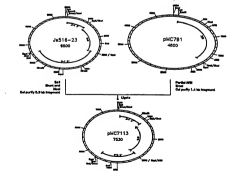
451 CCCGACTCCC TGCACCTGCC ATGG

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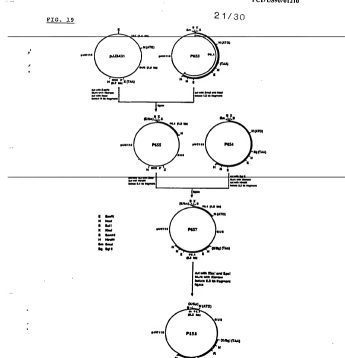
ě.	y pTDS133 (-468) y pTDS134 (-454)
, 1	AGGAATTECT CTCCATGGAT CCCCTCTATT TACCTGGCCA CCAAACATCC
^ 51	
101	AAATCTAAAA TAGTATACTT TAGCAGCCTC TCAATCTGAT TTGTTCCCCA
151	
201	CAGGGGGAGA GCCTAATCTT TCACGACTTT GTTTGTAACT GTTAGCCAGA
251	CCGGCGTATT TGTCAATGTA TARACACGTA ATARARAMMIA
301	AGTAAGACTT TGTATATAAG ACGTCACCTC TTAUGTGCAT GGTTATATACC
	GACATGTGCA GTGACGTTAT CAGATATAGC TCACCCTATA TATATAGCTC
401	TGTCCGGTGT CAGTGACAAT CACCATTCAT CAGCACCCCG GCAGTGCCAC

Service State Stat

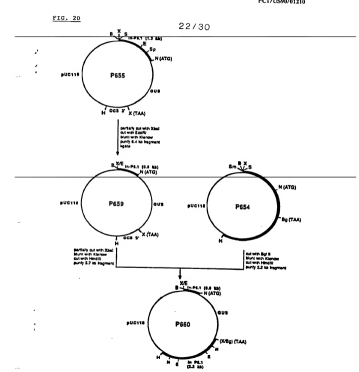
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FIG. 21

HPH443

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1 GAATTCTACG TACCATATAG TAAGACTTTG TATATAAGAC GTCACCTCTT

51 ACGTGCATGG TTATATGCGA CATGTGCAGT GACGTTAACC GCACCCTCCT

101 TCCCGTCGTT TCCCATCTCT TCCTCCTTTA GAGCTACCAC TATATAAATC

151 AGGGCTCATT TTCTCGCTCC TCACAGGCTC ATCAGCACCC CGGCAGTGCC
201 ACCCCGACTC CCTGCACCTG CCATGGCTT GGCTCGAGGT ACC

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FIG.22

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HPH 463

1 CTGCAGTACG TACCATATAG TAAGACTTTG TATATAAGAC GTCACCTCTT

51 ACGTGCATGG TATATAGGA CATGTGCAGT GACGTTATCA GATATAGGTC

101 ACCCTATATA TATAGCTCTG TCCGGTGTCA GTGACAATCA CCATTCATCT

151 CGCTTTGGAT CGATTGGTTT CGTAACTGGT GAAGGACTGA GGGTCTCGGA

201 GTGGATGATT TGGGGATTCTG TTCGAAGATT TGCGGAGGGG GGCCATGGCG

251 ACGGTACC

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HPH4 78

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1 GGATCCCCCG TACCATATGT AMGACTTTGT ATATAMGACG TCACCTCTTA

51 CGTGGATGGT TATATGGGAC ATGTCCAGTG ACGTTAACAA GGATCGGCGC

101 GCCACGCCGA GCTCGCCGGT ATATTTATAT TTGCTCAATG GACAGGCATG

151 GGGCTATCC GCTTTGGATC GATTGGTTC GTAACTGGTG AAGGACTGAG

201 GGTCTCGGAG TGGATGATTT GGGATTCTC TCGAAGATTT GCGGAGGGGG

251 GCCATGGCGA CGGTACC

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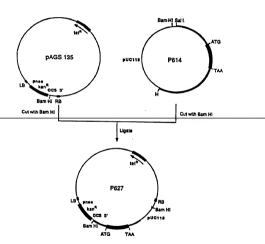
FIG. 24

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HPH 420

1	GAATTCTACG	TACCATATAG	TAAGACTTTG	TATATAAGAC	GTCACCTCTT
51	ACGTGCATGG	TTATATGCGA	CATGTGCAGT	GACGTTAACC	GCACCCTCCT
101	TCCCGTCGTT	TCCCATCTCT	TCCTCCTTTA	GAGCTACCAC	TATATAAAA
151	AGGGCTCATT	TTCTCGCTCC	TCACAGGCTC	ATCTCGCTTT	GGATCGATTG

201 GTTTCGTAAC TGGTGAAGGA CTGAGGGTCT CGGAGTGGAT GATTTGGGAT
251 TCTGTTCGAA GATTTGCGGA GGGGGGCCA<u>T</u> GGCGACGGTA CC



■ LB - TDNA left border

■ RB - TDNA right border

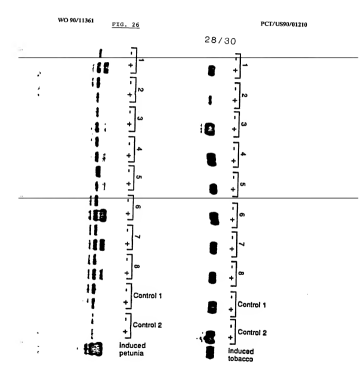
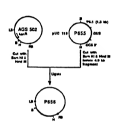
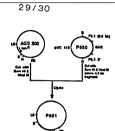
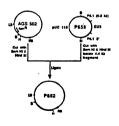
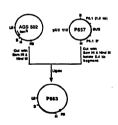


FIG. 27

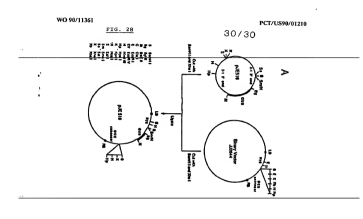


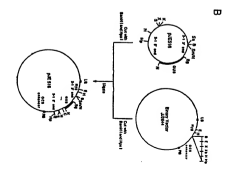






· IS - TONA left barrier · RS - TONA parts barrier





INTERNATIONAL SEARCH REPORT

International Application No PCT/US 90/01210

	SSIFICATION OF SUBJECT MATTER (il several classification (IPC) or to both National Patent Classification (IPC) or to both Nation					
IPC						
II. FIE	DS SEARCHED					
l	Minimum Documen					
Ciasini	Classification System Classification Symbols					
IPC ⁵	IPC ⁵ C 12 N 15/82, C 12 N 15/29, C 12 N 15/67 Documentation Searched other than Minimum Documentation to the Estant Intel surch Document are Included in the Patiest Searched					
III. DC	CUMENTS CONSIDERED TO BE RELEVANT	regrists of the relevant passages 12	Relayant to Claim No. 13			
Catagor	CHERRY OF DECUMENT, 17 WITH INDICATION, WITH SEPT	COLLEC, OF CHARLES				
P,	EP, A, 0332104 (CIBA-GEI 13 September 1989 see the whole docume		1-11,44-52			
A	Proceedings of the 1985 Protection Society C	British Crop	1-11			
	of herbicide antidot 1147-1153 see the whole articl page 1153 cited in the application	le, in particular				
-A- -E- -L- -D- -P-	* Special categories all citied documents: 'N **A document enhance the general state of the art which is not a considerable to the state of the art which is not citied to expect the state of the stat					
	May 1990	1 0. 07, 90				
1525						

	III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)				
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*	A	Plant Molecular Biology, vol. 7, 1986, Martinus Nijhoff Publishers, (Dordrecht, NL), R.C. Wiegand et al.: "Messenger RNA encoding a glutathione-S-transferase responsible for herbicide tolerance in maize is transferance in maize is transferance in page 236, column 1; page 241, column 2 - page 242, table 1 cited in the application EF, A, 0159884 (LUBRIZOL GENETICS INC.)	1,6-11	1 1	
	A	30 October 1985 see abstract; page 6, lines 1-7; page 9, lines 16-33; page 16, examples; claims cited in the application Nature, vol. 335, 29 September 1988, W.R. Marcotte Jr et al.: "Regulation	1,6-8,44-52		
ura.		of a wheat promoter by absolcie acid in rice protoplasts, pages 545-417 see the whole article cited in the application	1		
	P,A	EP, A, 0337532 (MOGEN INTERNATIONAL) 18 October 1989 see abstract; column 3, line 13 - column 5, line 35; column 5, line 51 - column 7, line 4; claims	1-11,44-52		
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9001210 SA 35373

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 27/06/90

The European Patent Office is in one way liable for these particulars which are merely given for the purpose of information.

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	10 09 09	AU-A-	3108089	14-09-89	
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